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L28

=> d stat que 795 SEA FILE=REGISTRY ABB=ON PLU=ON TETRAHYDROBORATE OR HYDROBORA L1 11 SEA FILE=REGISTRY ABB=ON PLU=ON SODIUM BOROHYDRATE?/CN OR L2 LITHIUM BOROHYDR?/CN OR POTASSIUM BOROHYDR?/CN L3 104214 SEA FILE=REGISTRY ABB=ON PLU=ON METAL OR METALS OR MAGNESIUM 128483 SEA FILE=REGISTRY ABB=ON PLU=ON BORATE L415523 SEA FILE=HCAPLUS ABB=ON PLU=ON L1 OR TETRAHYDROBORATE OR L5 HYDROBORATE L6 52 TERMS SEL PLU=ON L2 1- CHEM : L715422 SEA FILE=HCAPLUS ABB=ON PLU=ON L6 15583 SEA FILE=HCAPLUS ABB=ON PLU=ON L7 OR (SODIUM OR LITHIUM OR L8 POTASSIUM) (2A) BOROHYDR? L9 4137887 SEA FILE=HCAPLUS ABB=ON PLU=ON L3 OR METAL OR MAGNESIUM L10 166954 SEA FILE=HCAPLUS ABB=ON PLU=ON L4 OR BORATE L12 15254 SEA FILE=HCAPLUS ABB=ON PLU=ON L10(L)L9 L1438473 SEA FILE=HCAPLUS ABB=ON PLU=ON L10(L)(RCT/RL OR RACT/RL) L15 250013 SEA FILE=HCAPLUS ABB=ON PLU=ON L9(L)(RCT/RL OR RACT/RL) L16 2579 SEA FILE=HCAPLUS ABB=ON PLU=ON L12 AND L14 1952 SEA FILE=HCAPLUS ABB=ON PLU=ON L16 AND L15 L17 L18 104576 SEA FILE=REGISTRY ABB=ON PLU=ON PALLADIUM/BI 208 SEA FILE=REGISTRY ABB=ON PLU=ON PT/MF L19 L21 513099 SEA FILE=HCAPLUS ABB=ON PLU=ON L18 OR L19 OR PALLADIUM OR PLATINUM OR PD OR PT L25 3041 SEA FILE=HCAPLUS ABB=ON PLU=ON (L5 OR L8) (L) (PREP?/RL OR PREP OR PREPARTION) L26 78 SEA FILE=HCAPLUS ABB=ON PLU=ON L17 AND L25 L27 11 SEA FILE=HCAPLUS ABB=ON PLU=ON L26 AND L21

9 SEA FILE=HCAPLUS ABB=ON PLU=ON L27 AND PD=<NOVEMBER 25, 2003

=> =>

=> d ibib abs hitstr 128 1-9

L28 ANSWER 1 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN 2002:475940 HCAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 138:26390 TITLE: Hydrogen generation using sodium borohydride solution and metal catalyst coated on metal oxide AUTHOR (S): Kojima, Yoshitsugu; Suzuki, Ken-ichirou; Fukumoto, Kazuhiro; Sasaki, Megumi; Yamamoto, Toshio; Kawai, Yasuaki; Hayashi, Hiroaki CORPORATE SOURCE: Toyota Central Research & Development Laboratories, Inc., Nagakute-cho, Aichi-gun, Aichi, 480-1192, Japan SOURCE: International Journal of Hydrogen Energy (2002 ), 27(10), 1029-1034 CODEN: IJHEDX; ISSN: 0360-3199 Elsevier Science Ltd. PUBLISHER: DOCUMENT TYPE: Journal LANGUAGE: English Sodium borohydride (NaBH4) reacted slowly with water to liberate 4 mol of hydrogen/mol of the compound at room temperature Hydrogen generation was accelerated by applying metal-metal oxide catalysts such as Pt -TiO2, Pt-CoO and Pt-LiCoO2. As the size of metal crystallites decreased and the amount increased, the hydrogen generation rate increased. The hydrogen generation rates using Pt and LiCoO2 were high compared with those using other metal and metal oxide, resp. It seemed that a key finding was that use of the supercrit. CO2 method produced a superior catalyst. Borohydride ion was stabilized in alkaline solution containing at least 5 weight% of NaOH. Alkaline stabilized solution of NaBH4 can be applied as a hydrogen source. We found that Pt-LiCoO2 worked as an excellent catalyst for releasing hydrogen from the stabilized NaBH4 solution 7440-05-3, Palladium, uses 7440-06-4, IT Platinum, uses RL: CAT (Catalyst use); USES (Uses) (hydrogen generation by reaction of sodium borohydride with water using metal catalyst coated on metal oxide) 7440-05-3 HCAPLUS RNPalladium (8CI, 9CI) (CA INDEX NAME) CN Pd 7440-06-4 HCAPLUS RN Platinum (8CI, 9CI) (CA INDEX NAME) CN Pt TT 16940-66-2, Sodium borohydride RL: CPS (Chemical process); PEP (Physical, engineering or chemical process); RCT (Reactant); PROC (Process); RACT (Reactant or (hydrogen generation by reaction of sodium borohydride with water using metal catalyst coated on metal oxide) RN16940-66-2 HCAPLUS Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

CN

● Na+

REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L28 ANSWER 2 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:568925 HCAPLUS

DOCUMENT NUMBER: 121:168925

TITLE: Homoscorpionate (Tris(pyrazolyl)borate) Ligands

Containing Tethered 3-Phenyl Groups

AUTHOR(S): Rheingold, Arnold L.; Ostrander, Robert L.; Haggerty,

Brian S.; Trofimenko, Swiatoslaw

CORPORATE SOURCE: Department of Chemistry, University of Delaware,

Newark, DE, 19716-2522, USA

SOURCE: Inorganic Chemistry (1994), 33(17), 3666-76

CODEN: INOCAJ; ISSN: 0020-1669

DOCUMENT TYPE: Journal LANGUAGE: English

Four new homoscorpionate ligands were prepared, 3 of them involving a 3-Ph substituent subject to steric control via tethering to the the 4-position of the pyrazole ring. The tethering prevents major departure from parallel alignment of the Ph and pyrazolyl rings. The choice of the tether (methylene or 1,2-ethylene) results in the Ph 6'-CH being either pulled away from the metal in the former case or thrust toward the metal in the latter. The effect of the 5-Me substituent on the bite of the ligand was also explored. The new ligands were hydrotris(2H-benz[g]-4,5-dihydroindazol-2-yl)borate (=Tpa), hydrotris(3-methyl-2H-benz[g]-4,5-dihydroindazol-2-yl)borate (=Tpa, Me), hydrotris(1,4-dihydroindeno[1,2-c]pyrazol-1-yl)borate (=Tpb), and hydrotris(3-phenyl-5-methylpyrazol-1-yl)borate (=TpPh,Me), each of which differed subtly in its coordinative behavior from the other 3. Complexes L2M, LMX, LRh(COD), LRh(CO)2,  $LPd(\eta 3\text{-methallyl})$ , and  $LMo(CO)2(\eta 3\text{-methallyl})$  were synthesized, and the structures of TpaRh(CO)2, Tpa,MeZnI, TpbTl, and TpPh,MeZnI were determined by x-ray crystallog. TpaRh(CO)2 crystallizes in the space group P.hivin.1, with a 8.574(2), b 20.113(6), c 20.188(6) Å,  $\alpha$ 61.68(2),  $\beta$  84.14(2),  $\gamma$  85.15(2)° for Z = 4. Tpa,MeZnI crystallizes in the space group P212121, with a 10.002(1), b 16.237(3), c 19.952(3) Å for Z = 4. TpbTl crystallizes in the space group P21/n, with a 11.363(2), b = 11.096(2) Å, c 21.010(4) Å,  $\beta$ 99.33(2)° for Z = 4. The Ph and pyrazolyl planes are essentially coplanar. TpPh,MeZnI crystallizes in the space group Pna21, with a 32.645(6), b 11.327(3), c 16.180(3) Å for Z = 8. Cone and wedge angles were calculated for the new ligands and compared with revised previously reported values.

IT 157410-24-7P 157410-25-8P 157410-26-9P 157441-22-0P

RN 157410-24-7 HCAPLUS

CN Palladium,  $[(1,2,3-\eta)-2-methyl-2-propenyl]$  [tris(4,5-dihydro-3-methyl-2H-benz[g]indazolato-N2)hydroborato(1-)-N1,N1']- (9CI) (CA INDEX NAME)

Me 
$$\frac{H2}{C}$$
  $\frac{Me}{C}$   $\frac{H2}{C}$   $\frac{Me}{C}$   $\frac{H2}{C}$   $\frac{H2}{$ 

RN 157410-25-8 HCAPLUS

CN Palladium, [(1,2,3-η)-2-methyl-2-propenyl][tris(2,4-dihydroindeno[1,2-c]pyrazolato-N2)hydroborato(1-)-N1,N1']- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ N & & N \\ \hline \end{array}$$

RN 157410-26-9 HCAPLUS

CN Palladium, [hydrotris(5-methyl-3-phenyl-1H-pyrazolato-N1)borato(1-)-N2,N2'][(1,2,3-η)-2-methyl-2-propenyl]- (9CI) (CA INDEX NAME)

RN 157441-22-0 HCAPLUS

CN Palladium, [(1,2,3-η)-2-methyl-2-propenyl][tris(4,5-dihydro-2H-benz[g]indazolato-N2)hydroborato(1-)-N1,N1',N1'']- (9CI) (CA INDEX NAME)

IT 13762-51-1, Potassium tetrahydroborate(1-)

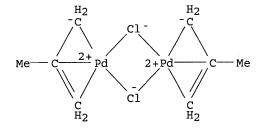
RL: RCT (Reactant); RACT (Reactant or reagent)
(reaction of, with benzdihydroindazoles or dihydroindenopyrazole or phenyl(methyl)pyrazole)

RN 13762-51-1 HCAPLUS

CN Borate(1-), tetrahydro-, potassium (8CI, 9CI) (CA INDEX NAME)

● K+

IT 12081-18-4, Bis(chloro( $\eta$ 3-2-methallyl) palladium) RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with thallous tris(pyrazolyl)borate derivs.) RN 12081-18-4 HCAPLUS CN Palladium, di- $\mu$ -chlorobis[(1,2,3- $\eta$ )-2-methyl-2-propenyl]di- (9CI) (CA INDEX NAME)



L28 ANSWER 3 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:437099 HCAPLUS

DOCUMENT NUMBER: 121:37099

TITLE: Palladium and aluminum thin film deposition

on thermally sensitive substrates from organometallic

complexes

AUTHOR (S): Datta, Saswati; Kim, Yoon Gi; Dowben, P. A.; Glass,

John A.; Kher, Shreyas S.; Peters, Scott A.; Spencer,

CORPORATE SOURCE: Int. Bus. Mach. Corp., Endicott, NY, 13760, USA

Metallized Plastics (1992), 3, 65-72 SOURCE:

CODEN: MPFAEU; ISSN: 1068-7440

DOCUMENT TYPE: Journal LANGUAGE: English

The UV decomposition of allylcyclopentadienyl palladium, [ $(\eta_3-C3H5)(\eta_5-C5H5)$  **Pd**] (I) has been used to deposit relatively pure (>95% palladium) metal films with submicrometer spatial resolution The films deposited on polyimide resins were found to be porous with no apparent damage to the substrate from the UV photoassisted decomposition technique. The fabrication of palladium thin films from I is consistent with the known energetics of the ligand to metal bonds in the complex. The room temperature metalization of thermally sensitive substrates, such as plastics, was also achieved from the decomposition of the volatile aluminum borohydride complex, AlH2(BH4).N(CH3)3. Films of

thicknesses between 100 Å and 2  $\mu m$  have been prepared and

characterized by SEM, Auger electron spectroscopy (AES), laser microprobe mass anal. (LAMMA), x-ray diffraction (XRD), and energy dispersive x-ray anal. (EDXA).

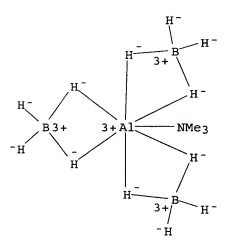
IT 19567-28-3, Aluminum tetrahydroborate, compound with NMe3 (1:1)

RL: RCT (Reactant); RACT (Reactant or reagent)

(decomposition of, chemical vacuum deposition of metal films on polyimides by)

19567-28-3 HCAPLUS RN

Aluminum, (N,N-dimethylmethanamine)tris[tetrahydroborato(1-)-H,H']-, CN (PB-7-12-22'2'2'')- (9CI) (CA INDEX NAME)



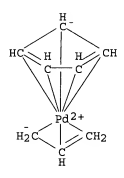
IT 1271-03-0, Allylcyclopentadienyl palladium

RL: RCT (Reactant); RACT (Reactant or reagent)

(decomposition of, laser radiation-induced, deposition of metal films on polyimides by)

1271-03-0 HCAPLUS RN

CN Palladium, (η5-2,4-cyclopentadien-1-yl) (η3-2-propenyl) - (9CI) (CA INDEX NAME)



IT 7440-05-3P, Palladium, preparation

RL: PREP (Preparation)

(films, deposition of, on polymeric substrates, by laser radiation-induced allylcyclopentadienyl palladium decomposition)

RN 7440-05-3 HCAPLUS

CN Palladium (8CI, 9CI) (CA INDEX NAME)

Pd

L28 ANSWER 4 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1990:525153 HCAPLUS

DOCUMENT NUMBER:

113:125153

TITLE:

Preparation of magnetic powder containing rare earth

elements

INVENTOR (S):

Buchkov, D.; Dragieva, I.; Iliev, I.; Todorov, O.;

Slavcheva-Staikova, M.

PATENT ASSIGNEE(S):

Higher Institute of Electrical and Mechanical

Engineering, Sofia, Bulg.

SOURCE:

Ger. (East), 3 pp.

CODEN: GEXXA8

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DD 271612	A7	19890913	DD 1986-291413	19860618 <
PRIORITY APPLN. INFO.:			DD 1986-291413	19860618

A solution of water-soluble salts of a rare earth metal, Fe, Ni, Co, Cu, and AB Pd at concns. of 1 + 10-3 to 1 + 10-1 mol/L is mixed with aqueous Na borohydride or Na hypophosphite (reducing agent) at 0.05-1 mol/L and 10-90° for 1-30 min in a magnetic field of (1-8) + 105 A/m to give a magnetic powder.

16940-66-2, Sodium borohydride IT

RL: RCT (Reactant); RACT (Reactant or reagent)

(reduction by, of metal salt solns., in preparation of magnetic powders)

RN16940-66-2 HCAPLUS

Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME) CN

Na +

L28 ANSWER 5 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1989:534235 HCAPLUS

DOCUMENT NUMBER: 111:134235

Unexpected regiochemistry in the reaction of TITLE:

benzotriazole with KBH4: synthesis of a new class of

poly(azolyl)borate ligands

AUTHOR (S): Lalor, F. J.; Miller, S.; Garvey, N. Dep. Chem., Univ. Coll., Cork, Ire. CORPORATE SOURCE:

Journal of Organometallic Chemistry (1988), SOURCE:

356(2), C57-60

CODEN: JORCAI; ISSN: 0022-328X

DOCUMENT TYPE: Journal LANGUAGE: English

CASREACT 111:134235 OTHER SOURCE(S):

The reaction of KBH4 with molten benzotriazole forms a series of poly(benzotriazolyl)borate anions [HnB(C6H4N3)4-n]- (n = 0-3) depending on the temperature The regiospecificity of the process differs from that of pyrazole/BH4- reactions in that B-N bond formation takes place in a manner that maximizes steric crowding at boron (i.e. at the triazole N(1) atoms). The complex-forming abilities of the new ligands have been investigated, and show some interesting differences from those of the

poly(1-pyrazolyl)borate analogs.

109782-45-8P 122627-92-3P 122658-64-4P IT

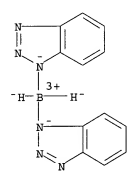
RL: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation); RACT (Reactant or reagent)

(preparation and reaction of, with transition metal complexes)

RN 109782-45-8 HCAPLUS

Borate(1-), bis(1H-benzotriazolato-κN1)dihydro-, potassium, (T-4)-CN (9CI) (CA INDEX NAME)



K+

122627-92-3 HCAPLUS RN

Borate(1-), tris(1H-benzotriazolato-κN1) hydro-, potassium, (T-4)-CN

(9CI) (CA INDEX NAME)

● K+

RN 122658-64-4 HCAPLUS

CN Borate(1-), tetrakis(1H-benzotriazolato-N1)-, potassium (9CI) (CA INDEX NAME)

• K+

IT 122651-40-5P 122651-42-7P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 122651-40-5 HCAPLUS

CN Palladium, (n3-2-propenyl) [tris(1H-benzotriazolato-N1)hydroborato(1-)-N2,N2']- (9CI) (CA INDEX NAME)

RN 122651-42-7 HCAPLUS

CN Palladium, (n3-2-propenyl) [tetrakis(1H-benzotriazolato-N1)borato(1-)-N2,N2']- (9CI) (CA INDEX NAME)

IT 13762-51-1, Potassium tetrahydroborate

RL: RCT (Reactant); RACT (Reactant or reagent)

(reaction of, with molten benzotriazole, regiochem. of)

RN 13762-51-1 HCAPLUS

CN Borate(1-), tetrahydro-, potassium (8CI, 9CI) (CA INDEX NAME)

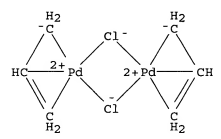
• K+

IT 12012-95-2

RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with poly(benzotriazolyl)borate anions)

RN 12012-95-2 HCAPLUS

CN Palladium,  $di-\mu$ -chlorobis ( $\eta 3-2$ -propenyl) di- (9CI) (CA INDEX NAME)



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L28 ANSWER 6 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN
                         1984:124631 HCAPLUS
ACCESSION NUMBER:
                         100:124631
DOCUMENT NUMBER:
                         Concentration of platinum metals by reducing
TITLE:
                         with sodium tetrahydroborate in the presence of
                         manganese dioxide
                         Khain, V. S.; Volkov, A. A.; Martynova, V. F.
AUTHOR (S):
CORPORATE SOURCE:
                         Ukhtinsk. Ind. Inst., Ukhta, USSR
                         Zhurnal Prikladnoi Khimii (Sankt-Peterburg, Russian
SOURCE:
                         Federation) (1984), 57(1), 30-3
                         CODEN: ZPKHAB; ISSN: 0044-4618
                         Journal
DOCUMENT TYPE:
LANGUAGE:
                         Russian
    Pt, Pd, and Rh were recovered from H2PtCl6, PdCl2, and
     RhCl3 solns. by reduction with NaBH4 at pH 4-10 in the presence of KMnO4.
     MnO2 formed by the reaction of KMnO4 with NaBH4 served as a collector for
     Pt-group metals.
IT
     7440-06-4P, preparation
     RL: PUR (Purification or recovery); PREP (Preparation)
        (recovery of, from hydrogen hexachloroplatinate solns., by reduction with
        sodium borohydride, manganese dioxide collector in)
RN
     7440-06-4 HCAPLUS
     Platinum (8CI, 9CI) (CA INDEX NAME)
CN
Ρt
     7440-05-3P, preparation
IT
     RL: PUR (Purification or recovery); PREP (Preparation)
        (recovery of, from palladium chloride solns., by reduction with
        sodium borohydride, manganese dioxide collector in)
RN
     7440-05-3 HCAPLUS
CN
     Palladium (8CI, 9CI) (CA INDEX NAME)
Pd
TΤ
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reduction by, of platinum-group metal chlorides, in
        aqueous solns.)
RN
     16940-66-2 HCAPLUS
     Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)
CN
```

IT 7647-10-1

● Na+

RL: RCT (Reactant); RACT (Reactant or reagent)
 (reduction of, with sodium borohydride in aqueous media, palladium
 recovery by)

RN 7647-10-1 HCAPLUS

CN Palladium chloride (PdCl2) (6CI, 8CI, 9CI) (CA INDEX NAME)

cl-Pd-Cl

```
L28 ANSWER 7 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN
                         1979:179411 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         90:179411
TITLE:
                         Reaction of sodium tetrahydroborate with precious
                         Mal'tseva, N. N.; Sterlyadkina, Z. K.; Erusalimchik,
AUTHOR (S):
                         I. G.; Mikheeva, V. I.
                         USSR
CORPORATE SOURCE:
SOURCE:
                         Zhurnal Neorganicheskoi Khimii (1979),
                         24(3), 822-4
CODEN: ZNOKAQ; ISSN: 0044-457X
DOCUMENT TYPE:
                         Journal
                         Russian
LANGUAGE:
     During the reaction of NaBH4 with aqueous solns. of Ag, Au, Pd, and
     Pt salts, highly pure free metal precipitated The stoichiometry of the
     oxidation-reduction reaction was established on the basis of the H evolved,
solution
     pH, and amount of reducing agent added. In acidic solution hydrolysis of NaBH4
     occurred simultaneously with the oxidation-reduction process. The
precipitated metal is
     highly dispersed and absorbs a significant amount of H.
     7440-05-3P, preparation
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, by reduction of palladium salts with sodium
        tetrahydroborate)
RN
     7440-05-3 HCAPLUS
CN
     Palladium (8CI, 9CI) (CA INDEX NAME)
Pd
IT
     7440-06-4P, preparation
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, by reduction of platinum salts with sodium
        tetrahydroborate)
RN
     7440-06-4 HCAPLUS
     Platinum (8CI, 9CI) (CA INDEX NAME)
CN
Pt
IT
     16940-66-2
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reduction by, of precious metal salts)
     16940-66-2 HCAPLUS
RN
     Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)
CN
```

## ● Na+

7440-05-3D, salts 7440-06-4D, salts 7440-22-4D IT , salts 7440-57-5D, salts
RL: RCT (Reactant); RACT (Reactant or reagent) (reduction of, by sodium tetrahydroborate)

7440-05-3 HCAPLUS RN

Palladium (8CI, 9CI) (CA INDEX NAME) CN

Pd

7440-06-4 HCAPLUS RN

Platinum (8CI, 9CI) (CA INDEX NAME) CN

Ρt

7440-22-4 HCAPLUS RN

Silver (8CI, 9CI) (CA INDEX NAME) CN

Ag

RN

7440-57-5 HCAPLUS Gold (8CI, 9CI) (CA INDEX NAME) CN

Au

L28 ANSWER 8 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1972:493257 HCAPLUS DOCUMENT NUMBER: 77:93257 TITLE: Catalysts formed by the reduction of Group VIII metal salts with sodium borohydride Sterlyadkina, Z. K.; Mal'tseva, N. N.; Frangulyan, G. AUTHOR (S): D.; Polkovnikov, B. D.; Bakulina, V. M. CORPORATE SOURCE: Inst. Org. Khim. im. Zelinskogo, Moscow, USSR Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya ( SOURCE: 1972), (6), 1240-5 CODEN: IASKA6; ISSN: 0002-3353 DOCUMENT TYPE: Journal Russian LANGUAGE: The ppts. formed by reduction of NiCl2 and CoCl2 by NaBH4 contain the free metal and M2B (M = Co, Ni) after being heated in Ar at 400°. Similar reduction of PdCl2 and H2PtCl6 leads to Pt and  $\alpha$ -7440-06-4P, uses and miscellaneous IT RL: CAT (Catalyst use); PREP (Preparation); USES (Uses) (catalysts, reduction of chloroplatinic acid by sodium borohydride in preparation of) 7440-06-4 HCAPLUS RN Platinum (8CI, 9CI) (CA INDEX NAME) CN Ρt TT 7440-05-3P, uses and miscellaneous RL: CAT (Catalyst use); PREP (Preparation); USES (Uses) (catalysts, reduction of palladium chloride by sodium borohydride in preparation of) RN7440-05-3 HCAPLUS Palladium (8CI, 9CI) (CA INDEX NAME) CNPd IT 16940-66-2 RL: RCT (Reactant); RACT (Reactant or reagent) (reduction by, of Group VIII metal salts in catalyst preparation) 16940-66-2 HCAPLUS RN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME) CN

Na +

L28 ANSWER 9 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1971:71098 HCAPLUS

DOCUMENT NUMBER: 74:71098

TITLE: Tertiary phosphine-hydride and -hydridoborohydride

compounds of nickel and palladium

AUTHOR(S): Green, Malcolm L. H.; Munakata, H.; Saito, Tato

CORPORATE SOURCE: Inorg. Chem. Lab., Oxford, UK

SOURCE: Journal of the Chemical Society [Section] A: Inorganic, Physical, Theoretical (1971),

(3), 469-74

CODEN: JCSIAP; ISSN: 0022-4944

DOCUMENT TYPE: Journal LANGUAGE: English

AB The isolation and properties of the complexes trans-(R13P)2M1H(BH4) and

trans-(R13P)2M2HX, where M1 = Ni or Pd; R1 = iso-Pr, cyclohexyl;

M2 = Pd; X = Cl, Br, I, or NCS, are described. Phosphine

exchange reactions of (R13P)2MHX with R23P, where R2 = Et, Pr, and Bu,

were studied by 1H NMR.

IT 16971-29-2DP, Borate(1-), tetrahydro-, Group VIII

metal complexes 27900-91-0P 28016-71-9P 28016-73-1P 30916-07-5P 31178-50-4P

31178-51-5P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) 16971-29-2 HCAPLUS

CN Borate(1-), tetrahydro- (8CI, 9CI) (CA INDEX NAME)

RN

RN 27900-91-0 HCAPLUS

CN Palladium, chlorohydrobis[tris(1-methylethyl)phosphine]-, (SP-4-3)- (9CI) (CA INDEX NAME)

RN 28016-71-9 HCAPLUS

\_\_\_\_\_\_

RN 28016-73-1 HCAPLUS

CN Palladium, bromohydrobis(tricyclohexylphosphine)-, trans- (8CI) (CA INDEX NAME)

RN 30916-07-5 HCAPLUS

CN Palladium, hydro[tetrahydroborate(1-)]bis(triisopropylphosphine)-, trans-(8CI) (CA INDEX NAME)

RN 31178-50-4 HCAPLUS

CN Palladium, hydroiodobis(tricyclohexylphosphine)-, trans- (8CI) (CA INDEX NAME)

RN 31178-51-5 HCAPLUS

CN Palladium, hydro(isothiocyanato)bis(tricyclohexylphosphine)-, trans- (8CI) (CA INDEX NAME)

IT 17523-47-6 24899-12-5

RL: RCT (Reactant); RACT (Reactant or reagent)
(redistribution reaction of, with Group VIII metal
trialkylphosphine complexes, NMR in relation to)

RN 17523-47-6 HCAPLUS

CN Palladium, dichlorobis(tributylphosphine)-, (SP-4-1)- (9CI) (CA INDEX NAME)

RN 24899-12-5 HCAPLUS

CN Nickel, hydro[tetrahydroborato(1-)-H,H']bis(tricyclohexylphosphine)-, (TB-5-11)- (9CI) (CA INDEX NAME)

IT 15642-19-0

RL: RCT (Reactant); RACT (Reactant or reagent)
(redistribution reaction of, with Group VIII metal
tricyclohexylphosphine complexes, NMR in relation to)

RN 15642-19-0 HCAPLUS

CN Palladium, dichlorobis(triethylphosphine)-, (SP-4-1)- (9CI) (CA INDEX NAME)

IT 17523-48-7

RL: RCT (Reactant); RACT (Reactant or reagent) (redistribution reaction of, with nickel tricyclohexylphosphine complexes, NMR in relation to)

RN 17523-48-7 HCAPLUS

CN Palladium, dibromobis(tributylphosphine)-, trans- (8CI) (CA INDEX NAME)

IT 30916-06-4

RN

RL: RCT (Reactant); RACT (Reactant or reagent)
 (redistribution reaction of, with palladium trialkyl
 phosphine complexes, NMR in relation to)
30916-06-4 HCAPLUS

IT 31095-23-5

RL: RCT (Reactant); RACT (Reactant or reagent)
(redistribution reaction of, with palladium tributyl
phosphine complexes, NMR in relation to)

RN 31095-23-5 HCAPLUS

CN Nickel, hydro[tetrahydroborato(1-)-H,H']bis[tris(1-methylethyl)phosphine]-, (TB-5-11)- (9CI) (CA INDEX NAME)

=> =>	d stat que	e 129
L1	795	SEA FILE=REGISTRY ABB=ON PLU=ON TETRAHYDROBORATE OR HYDROBORA
		TE
L2	11	SEA FILE=REGISTRY ABB=ON PLU=ON SODIUM BOROHYDRATE?/CN OR
		LITHIUM BOROHYDR?/CN OR POTASSIUM BOROHYDR?/CN
L3	104214	SEA FILE=REGISTRY ABB=ON PLU=ON METAL OR METALS OR MAGNESIUM
L4		SEA FILE=REGISTRY ABB=ON PLU=ON BORATE
L5	15523	SEA FILE=HCAPLUS ABB=ON PLU=ON L1 OR TETRAHYDROBORATE OR
		HYDROBORATE
L6		SEL PLU=ON L2 1- CHEM : 52 TERMS
L7		SEA FILE=HCAPLUS ABB=ON PLU=ON L6
L8	15583	SEA FILE=HCAPLUS ABB=ON PLU=ON L7 OR (SODIUM OR LITHIUM OR
		POTASSIUM) (2A) BOROHYDR?
L9		SEA FILE=HCAPLUS ABB=ON PLU=ON L3 OR METAL OR MAGNESIUM
L10		SEA FILE=HCAPLUS ABB=ON PLU=ON L4 OR BORATE
L12		SEA FILE=HCAPLUS ABB=ON PLU=ON L10(L)L9
L14		SEA FILE=HCAPLUS ABB=ON PLU=ON L10(L)(RCT/RL OR RACT/RL)
L15		SEA FILE=HCAPLUS ABB=ON PLU=ON L9(L)(RCT/RL OR RACT/RL)
L16		SEA FILE=HCAPLUS ABB=ON PLU=ON L12 AND L14
L17		SEA FILE=HCAPLUS ABB=ON PLU=ON L16 AND L15
L18		SEA FILE=REGISTRY ABB=ON PLU=ON PALLADIUM/BI SEA FILE=REGISTRY ABB=ON PLU=ON PT/MF
L19		SEA FILE=REGISTRY ABB=ON PLU=ON P1/MF SEA FILE=HCAPLUS ABB=ON PLU=ON L18 OR L19 OR PALLADIUM OR
L21	513099	PLATINUM OR PD OR PT
L25	2041	SEA FILE=HCAPLUS ABB=ON PLU=ON (L5 OR L8)(L)(PREP?/RL OR
ь25	3041	PREP OR PREPARTION)
L26	70	SEA FILE=HCAPLUS ABB=ON PLU=ON L17 AND L25
L27		SEA FILE=HCAPLUS ABB=ON PLU=ON L26 AND L21
L28		SEA FILE=HCAPLUS ABB=ON PLU=ON L27 AND PD= <november 2003<="" 25,="" td=""></november>
1120	9	DEA I I DEL MONTE DO DESCON PER DE LA PROPRIO DE CONTRA
L29	2	SEA FILE=HCAPLUS ABB=ON PLU=ON L27 NOT L28

<sup>=&</sup>gt; d ibib abs hitstr 129 1-2

L29 ANSWER 1 OF 2 HCAPLUS COPYRIGHT 2006 ACS on STN

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ACCESSION NUMBER:
                         2004:676920 HCAPLUS
DOCUMENT NUMBER:
                         142:6582
TITLE:
                         Hydrocarbon C-H bond activation with Tp'Pt
                         complexes
AUTHOR (S):
                         Norris, Cynthia M.; Templeton, Joseph L.
CORPORATE SOURCE:
                         Department of Chemistry, University of North Carolina,
                         Chapel Hill, NC, 27599-3290, USA
SOURCE:
                         ACS Symposium Series (2004), 885 (Activation and
                         Functionalization of C-H Bonds), 303-318
                         CODEN: ACSMC8; ISSN: 0097-6156
PUBLISHER:
                         American Chemical Society
DOCUMENT TYPE:
                         Journal; General Review
                         English
LANGUAGE:
    A review. Stabilization of Pt complexes resembling key
     intermediates in C-H bond activation has been achieved through the use of
     the strongly electron-donating hydridotris(3,5-dimethyl-1H-pyrazol-1-
     yl)borate (Tp') ligand. Isolation of stable alkyl(hydrido)
     platinum(IV) complexes, Tp'PtMe2H and Tp'PtMeH2, allowed the
     authors to study the mechanism of reductive alkane elimination from these
     complexes via (1) thermolysis, (2) Lewis acid addition, and (3) low temperature
     protonation. By replacing the alkyl group with SiR3 and C6R5 in the low
     temperature protonation of Tp'PtRH2, isolation of five-coordinate Pt
     (IV) complexes and Pt(II) η2-arene adducts, resp., was
     achieved. These Pt(II)/Pt(IV) interconversions
     provide insight into the mechanisms by which these reagents activate
     strong C-H bonds and also provide a foundation for future plans to
     functionalize hydrocarbons.
IT
     7440-06-4DP, Platinum, hydridotris(3,5-dimethylpyrazol-1-
     yl)borate complexes 83534-02-5DP, Hydridotris (3,5-
     dimethylpyrazol-1-yl)borate, platinum complexes
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); PROC (Process); RACT (Reactant or reagent)
        (hydrocarbon carbon-hydrogen bond activation with platinum
        hydridotripyrazolylborato complexes)
     7440-06-4 HCAPLUS
RN
     Platinum (8CI, 9CI) (CA INDEX NAME)
CN
Pt
RN
     83534-02-5 HCAPLUS
CN
     Borate(1-), tris(3,5-dimethyl-1H-pyrazolato-\kappaN1)hydro-, (T-4)- (9CI)
       (CA INDEX NAME)
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$$\begin{array}{c} \text{Me} \\ \text{N} \\ \text{N} \\ \text{Me} \\ \text{N} \\ \text{N} \\ \text{Me} \\ \text{Me} \\ \\ \text{Me} \\ \end{array}$$

REFERENCE COUNT:

91 THERE ARE 91 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Valenrod 10.721479 L29 ANSWER 2 OF 2 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:651206 HCAPLUS DOCUMENT NUMBER: 141:192634 TITLE: method to produce tetrahydroborate salts INVENTOR (S): Suda, Seijiro; Li, Chou-Peng; Iwase, Yasuyoshi; Morigasaki, Nobuto PATENT ASSIGNEE(S): Materials & Energy Research Institute Tokyo, Ltd. MERIT., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 9 pp. CODEN: JKXXAF DOCUMENT TYPE: Patent Japanese LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE ---------\_\_\_\_\_ -----JP 2004224593 A2 20040812 JP 2003-11299 20030120 PRIORITY APPLN. INFO.: JP 2003-11299 20030120 The method includes hydriding reaction of alkali or alkaline earth metal meta-borate by supplying mixed gas of O2-accepting gas (e.g. CO, C≤5 hydrocarbons), H2, and O2 with a hydriding catalyst selected from ≥1 of Ni, Co, Fe, Pt, Cu, Pd, Ru, and Rh. The borate raw material is obtained from spent fuel cells. The method enhances the hydriding reaction under mild conditions, and produces NaBH4, LiBH4, and KBH4 in low operation costs. TT 14100-65-3, Meta-borate RL: RCT (Reactant); RACT (Reactant or reagent) (alkali metal salt, alkaline metal salt; method to produce tetrahydroborate salts) 14100-65-3 HCAPLUS RNCN Borate (BO21-) (8CI, 9CI) (CA INDEX NAME) -o- B= o 7440-05-3, Palladium, uses 7440-06-4, TT Platinum, uses RL: CAT (Catalyst use); USES (Uses) (method to produce tetrahydroborate salts) 7440-05-3 HCAPLUS RN Palladium (8CI, 9CI) (CA INDEX NAME) CN Pd 7440-06-4 HCAPLUS RNCN Platinum (8CI, 9CI) (CA INDEX NAME) Pt

IT 13762-51-1P, Potassium Tetrahydroborate 16940-66-2P, Sodium Tetrahydroborate 16949-15-8P, Lithium Tetrahydroborate RL: PUR (Purification or recovery); PREP (Preparation)

(method to produce tetrahydroborate salts) RN

13762-51-1 HCAPLUS

Borate(1-), tetrahydro-, potassium (8CI, 9CI) (CA INDEX NAME)

CN

● K+

16940-66-2 HCAPLUS Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME) CN

• Na+

RN16949-15-8 HCAPLUS Borate(1-), tetrahydro-, lithium (8CI, 9CI) (CA INDEX NAME)

• Li+

=> => d	l stat que	e
L1		SEA FILE=REGISTRY ABB=ON PLU=ON TETRAHYDROBORATE OR HYDROBORA TE
L2	11	SEA FILE=REGISTRY ABB=ON PLU=ON SODIUM BOROHYDRATE?/CN OR
		LITHIUM BOROHYDR?/CN OR POTASSIUM BOROHYDR?/CN
L3	104214	SEA FILE=REGISTRY ABB=ON PLU=ON METAL OR METALS OR MAGNESIUM
L4	128483	SEA FILE=REGISTRY ABB=ON PLU=ON BORATE
L5	15523	SEA FILE=HCAPLUS ABB=ON PLU=ON L1 OR TETRAHYDROBORATE OR
		HYDROBORATE
L6		SEL PLU=ON L2 1- CHEM : 52 TERMS
L7	15422	SEA FILE=HCAPLUS ABB=ON PLU=ON L6
L8	15583	SEA FILE=HCAPLUS ABB=ON PLU=ON L7 OR (SODIUM OR LITHIUM OR
		POTASSIUM) (2A) BOROHYDR?
L9		SEA FILE=HCAPLUS ABB=ON PLU=ON L3 OR METAL OR MAGNESIUM
L10		SEA FILE=HCAPLUS ABB=ON PLU=ON L4 OR BORATE
L12		SEA FILE=HCAPLUS ABB=ON PLU=ON L10(L)L9
L14		SEA FILE=HCAPLUS ABB=ON PLU=ON L10(L)(RCT/RL OR RACT/RL)
L15		SEA FILE=HCAPLUS ABB=ON PLU=ON L9(L) (RCT/RL OR RACT/RL)
L16		SEA FILE=HCAPLUS ABB=ON PLU=ON L12 AND L14
L17		SEA FILE=HCAPLUS ABB=ON PLU=ON L16 AND L15
L18		SEA FILE=REGISTRY ABB=ON PLU=ON PALLADIUM/BI
L19		SEA FILE=REGISTRY ABB=ON PLU=ON PT/MF
L21	513099	SEA FILE=HCAPLUS ABB=ON PLU=ON L18 OR L19 OR PALLADIUM OR
		PLATINUM OR PD OR PT
L25	3041	SEA FILE=HCAPLUS ABB=ON PLU=ON (L5 OR L8)(L)(PREP?/RL OR
		PREP OR PREPARTION)
L26		SEA FILE=HCAPLUS ABB=ON PLU=ON L17 AND L25
L27		SEA FILE=HCAPLUS ABB=ON PLU=ON L26 AND L21
L28	9	SEA FILE=HCAPLUS ABB=ON PLU=ON L27 AND PD= <november 2003<="" 25,="" td=""></november>
L30	524	SEA FILE=HCAPLUS ABB=ON PLU=ON (L5 OR L8)(L)(PURIFI?/RL OR
		PURIFI? OR RECOVER?)
L31		SEA FILE=HCAPLUS ABB=ON PLU=ON L30 AND L10 AND L9
L32		SEA FILE=HCAPLUS ABB=ON PLU=ON L31 AND L9 AND L21
L33	36	SEA FILE=HCAPLUS ABB=ON PLU=ON L32 NOT (L27 OR L28)

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=> d ibib abs hitstr 133 1-36

L33 ANSWER 1 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:652766 HCAPLUS

DOCUMENT NUMBER: 144:153746

TITLE: Recovery of platinum from spent catalysts

from catalytic reforming unit using the Gemini process AUTHOR(S): Soltan-Mohammadzadeh, Jafar-Sadeg; Fatehifar, Esmaeel;

Hamedi, Hassan

CORPORATE SOURCE: Chem. Engineering, Sahand Technical University,

Tabriz, Iran

SOURCE: Iranian National Chemical Engineering Congress, 8th,

Mashhad, Islamic Republic of Iran, Oct. 19-21, 2003 (2003), 227/1-227/11. Danishgah-i Ferdowsi Mashhad,

Danishkada Muhandisi: Mashhad, Iran.

CODEN: 69GMYP

DOCUMENT TYPE: Conference; (computer optical disk)

LANGUAGE: Persian

AB Studies were carried out to optimize the conditions for the solubilization of base catalyst alumina and concentration and separation of Pt using formic

acid and NaBH4. Pt recovery amounted to 94.1% with formic acid

and 95.68% with NaBH4.

IT 16940-66-2, Sodium borohydride

RL: NUU (Other use, unclassified); USES (Uses)
(recovery of platinum from spent catalysts from catalytic reforming unit using Gemini process)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

• Na+

IT 7440-06-4P, Platinum, preparation

RL: PUR (Purification or recovery); PREP (Preparation) (recovery of platinum from spent catalysts from catalytic reforming unit using Gemini process)

RN 7440-06-4 HCAPLUS

CN Platinum (8CI, 9CI) (CA INDEX NAME)

Ρt

L33 ANSWER 2 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:395865 HCAPLUS

DOCUMENT NUMBER: 142:396971

TITLE: Process for removal of heavy metal ions from wastewater using reducing agent composition

INVENTOR(S): Covaliov, Victor; Covaliova, Olga; Duca, Gheorghe

PATENT ASSIGNEE(S): Universitatea de Stat din Moldova, Moldova

SOURCE: Mold. Unexam. Pat. Appl., 7 pp.

CODEN: MDXXAX

DOCUMENT TYPE: Patent LANGUAGE: Romanian

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

MD 990123 A 20001130 MD 1999-123 19990317

MD 1649 F2 20010430

PRIORITY APPLN. INFO.: MD 1999-123 19990317

The invention refers to processes for purification of sewage from. Heavy metal ions are removed from wastewater using a chemical reducing agent composition containing sodium borohydride

0.05-0.06, sodium hypophosphite 0.25-0.30, formaldehyde 0.60-0.70, and hydrazine 0.30-0.40 mol/L at pH 6-8. The process is realized in a flow at the linear velocity of 0.2-0.3 m/min in an electromagnetic field at a voltage of 20-30 W/dm3and frequency of 60-74 kHz on the catalyst surface. A nickel-zinc alloy is used as catalyst and is coated by electrochem. deposition onto a steel grill with the cell dimension of 0.5-2.0 mm. The solution is added so that the weight ratio to the total heavy metal ions content is (1.5-2.0):1. Use of this composition results in increased sewage purification degree and decreased material and power consumption for the purification process.

IT 7440-02-0, Nickel, processes 7440-05-3,

Palladium, processes 7440-22-4, Silver, processes

7440-50-8, Copper, processes 7440-66-6, Zinc, processes

RL: REM (Removal or disposal); PROC (Process)

(removal of heavy metal ions from wastewater using reducing agent composition)

RN 7440-02-0 HCAPLUS

CN Nickel (8CI, 9CI) (CA INDEX NAME)

Ni

RN 7440-05-3 HCAPLUS

CN Palladium (8CI, 9CI) (CA INDEX NAME)

Pd

RN 7440-22-4 HCAPLUS

CN Silver (8CI, 9CI) (CA INDEX NAME)

Αg

RN 7440-50-8 HCAPLUS

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Valenrod 10_721479
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CNCopper (7CI, 8CI, 9CI) (CA INDEX NAME)

Cu

7440-66-6 HCAPLUS Zinc (7CI, 8CI, 9CI) (CA INDEX NAME) RNCN

Zn

16940-66-2, Sodium borohydride ITRL: NUU (Other use, unclassified); USES (Uses) (removal of heavy metal ions from wastewater using reducing agent composition containing)

RN 16940-66-2 HCAPLUS CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

• Na+

L33 ANSWER 3 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN 2004:780106 HCAPLUS ACCESSION NUMBER: 141:245652 DOCUMENT NUMBER: TITLE: Production and purification of hydrogen INVENTOR (S): Ring, Terry A.; Freise, William; Maylett, Brett; Fisher, Matthew PATENT ASSIGNEE(S): USA U.S. Pat. Appl. Publ., 8 pp. SOURCE: CODEN: USXXCO DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION: KIND DATE APPLICATION NO. DATE PATENT NO. --------------US 2003-384961 -----US 2004184987 A1 20040923 20030310 PRIORITY APPLN. INFO.: Highly pure hydrogen gas is produced by reaction of Na, NaH, NaAlH4, NaBH4, Li, LiH, or LiBH4 with water and subsequent purification of the hydrogen using a hydrogen-selective **metal** membrane. The membrane can consist of **Pd**, Ta, V, Nb, Y, Th, Zr, or Ti. The membrane is supported by a rigid porous structural support. 7440-05-3, Palladium, uses 7440-32-6, IT Titanium, uses RL: DEV (Device component use); USES (Uses) (membrane; production and purification of hydrogen) 7440-05-3 HCAPLUS RNPalladium (8CI, 9CI) (CA INDEX NAME) CNPd RN 7440-32-6 HCAPLUS Titanium (8CI, 9CI) (CA INDEX NAME) CN Τi TΤ 7440-23-5, Sodium, reactions 16940-66-2, Sodium borohydride (NaBH4) 16949-15-8, Lithium borohydride (LiBH4) RL: CPS (Chemical process); PEP (Physical, engineering or chemical process); RCT (Reactant); PROC (Process); RACT (Reactant or reagent) (production and purification of hydrogen) 7440-23-5 HCAPLUS RN Sodium (8CI, 9CI) (CA INDEX NAME) CN Na 16940-66-2 HCAPLUS RN

Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

CN

• Na+

RN 16949-15-8 HCAPLUS CN Borate(1-), tetrahydro-, lithium (8CI, 9CI) (CA INDEX NAME)

● Li+

L33 ANSWER 4 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:112383 HCAPLUS

DOCUMENT NUMBER: 140:321195

TITLE: Process Development of a Scaleable Route to

(2R)-[3-(2-Aminopropyl)-1H-indol-7-yloxy]-N,N-diethylacetamide: A Key Intermediate for AJ-9677, a Potent and Selective Human and Rat  $\beta$ 3-Adrenergic

Receptor Agonist

AUTHOR(S): Harada, Hiroshi; Fujii, Akihito; Odai, Osamu; Kato,

Shiro

CORPORATE SOURCE: Chemistry Research Laboratories, Dainippon

Pharmaceutical Co., Ltd., Suita, Osaka, 564-0053,

Japan

SOURCE: Organic Process Research & Development (2004), 8(2),

Ι

238-245

CODEN: OPRDFK; ISSN: 1083-6160

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 140:321195

GI

$$Et_2N$$
 $NH_2$ 
 $Me$ 
 $NH_2$ 

$$\begin{array}{c|c} & \text{OH} & \text{OH} \\ & \text{H} & \text{N} & \text{Cl} \\ & \text{HO}_2\text{C} & \text{H} & \text{II} \end{array}$$

AB Nonracemic (aminopropylindoleoxy) acetamide I, an intermediate in the preparation of β3-adrenergic receptor agonist AJ-9677 II, is prepared in five steps and 43% overall yield from 7-benzyloxy-1H-indole without chromatog. purification The key step in the preparation of I is a regioselective and stereoselective acylation of 7-benzyloxy-1H-indole with N-Fmoc-D-alaninyl chloride; reduction of the acylindole intermediate with

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sodium borohydride in a mixture of acetonitrile and
     isopropanol, and salt formation yields the nonracemic 1:1 oxalate salt of
     III. Protection of the terminal amine with Boc anhydride, a one-step
     debenzylation and alkylation using hydrogenation with palladium
    on carbon followed by alkylation of the phenol with N,N-di-Et
     chloroacetamide, removal of the Boc group with oxalic acid and treatment
    of the crystalline oxalate salt with potassium carbonate yields I.
     Deprotonation of 7-benzyloxy-1H-indole using methylmagnesium bromide is
    more effective than deprotonation with other Me Grignard reagents.
    N-Fmoc-D-alaninyl chloride is a more effective reagent in the preparation of I
    than other N-protected D-alaninyl chlorides; N-Cbz-D-alaninyl chloride
    does not react with the anion generated from 7-benzyloxy-1H-indole and
    methylmagnesium bromide, a phthaloyl-protected alaninyl chloride reacts
    but yields an acylindole which could not be reduced, and
    N-trifluoroacetyl-D-alaninyl chloride gives a lower yield in the acylation
     step. Reduction with sodium borohydride gives better
     results than other reducing agents such as lithium
    borohydride, borane-THF, and Vitride in the reduction of the
     acylindole intermediate in the preparation of I; using a mixed solvent system
     of acetonitrile and isopropanol gives better results than the use of
     either an alc. solvent alone or the use of other alcs. such as ethanol as
     cosolvents with acetonitrile. Oxalic acid salts of the aminopropylindole
     and the aminopropylindoleoxyacetamide intermediates in the preparation of I are
     crystalline, while salts with other inorg. and organic acids tried are
amorphous.
    Alkylation of the penultimate phenol intermediate with chloroacetic acid
     esters gives lactam products rather than the desired
     aminopropylindoleoxyacetate esters.
     676-58-4, Methylmagnesium chloride 917-64-6,
     Methylmagnesium iodide
     RL: RGT (Reagent); RACT (Reactant or reagent)
        (less effective methylmagnesium halide reagent for the deprotonation of
        7-benzyloxyindole in the regioselective acylation with N-protected
        D-alaninyl chlorides for the synthesis of the \beta3-adrenergic
        receptor agonist AJ-9677)
     676-58-4 HCAPLUS
     Magnesium, chloromethyl- (7CI, 8CI, 9CI) (CA INDEX NAME)
H_3C-Mg-Cl
     917-64-6 HCAPLUS
     Magnesium, iodomethyl- (7CI, 8CI, 9CI) (CA INDEX NAME)
H_3C-Mg-I
     16949-15-8, Lithium borohydride
     RL: RGT (Reagent); RACT (Reactant or reagent)
        (less effective reducing agent tried in the reduction of a nonracemic
        3-acylindole in the preparation of an (aminopropylindoleoxy)acetamide
        intermediate for the synthesis of the \beta3-adrenergic receptor
        agonist AJ-9677)
     16949-15-8 HCAPLUS
     Borate(1-), tetrahydro-, lithium (8CI, 9CI) (CA INDEX NAME)
```

TT

RN

RN

CN

TΤ

ВN

CN

• Li+

TT 75-16-1, Methylmagnesium bromide
RL: RGT (Reagent); RACT (Reactant or reagent)
 (optimal methylmagnesium halide reagent for the deprotonation of
 7-benzyloxyindole in the regioselective acylation with N-protected
 D-alaninyl chlorides for the synthesis of the β3-adrenergic
 receptor agonist AJ-9677)

RN 75-16-1 HCAPLUS

CN Magnesium, bromomethyl- (8CI, 9CI) (CA INDEX NAME)

Br-Mg-CH3

IT 16940-66-2, Sodium borohydride

RL: RGT (Reagent); RACT (Reactant or reagent) (optimal reducing agent tried in the reduction of a nonracemic 3-acylindole in the preparation of an (aminopropylindoleoxy) acetamide intermediate for the synthesis of the  $\beta$ 3-adrenergic receptor agonist AJ-9677)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

● Na+

REFERENCE COUNT: 19 THERE ARE 19 CIT

19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L33 ANSWER 5 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:767751 HCAPLUS

DOCUMENT NUMBER: 139:276637

TITLE: Purification of ethylenic compounds having fluorinated

organic group

INVENTOR(S): Matsuda, Takashi; Koike, Noriyuki; Oyama, Masayuki

PATENT ASSIGNEE(S): Shin-Etsu Chemical Industry Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

OTHER SOURCE(S): MARPAT 139:276637

AB Rf(CH:CH2)n (Rf= mono- or divalent F-containing organic group; n = 1, 2) are purified by treating with ≥1 selected from metal hydrides and metal-hydrogen complex compds. to eliminate I from iodine compds. contained as impurities and other components which interfere Pt-catalyzed hydrosilylation reaction. A mixture of C4F9CH;CH2 (I; 100 g, I2 content 105 ppm), NaBH4, NaOH solution, and Me2CHOH was stirred ay 60° for 1 h to give 97 g I containing ≤1 ppm I2. The purified I was treated with MeSiHCl2 in Me2CHOH containing H2PtCl6 at 150° for 18 h to show conversion 98.2%.

IT 16940-66-2, Sodium borohydride

RL: RGT (Reagent); RACT (Reactant or reagent)
 (purification of F-containing ethylenic compds. by eliminating I from
 impurity iodides using metal hydrides or metal
 -hydrogen complexes)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

Na +

L33 ANSWER 6 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2003:673831 HCAPLUS DOCUMENT NUMBER: 139:179794 TITLE: Method for purification of methylal containing peroxides INVENTOR(S): Omori, Hideki; Shoji, Hiroshi; Nakamura, Tomizo PATENT ASSIGNEE(S): Maruzen Oil Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 4 pp. CODEN: JKXXAF DOCUMENT TYPE: Patent Japanese LANGUAGE: FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE -------------------------A2 JP 2002-34153 JP 2002-34153 JP 2003238471 20030827 20020212 PRIORITY APPLN. INFO.: 20020212 The method includes reduction of methylal (I) containing peroxides and adsorption of the reduced products to remove HCO2H (II). Thus, I containing 180 ppm peroxide was hydrogenolyzed over Raney Ni to give a product (peroxide content <2 ppm, II content 60 ppm), which was treated with Mol. Sieve 4A, showing II content <1 ppm. 1318-93-0, Montmorillonite, uses IT RL: NUU (Other use, unclassified); USES (Uses) (adsorbent; purification of methylal containing peroxides by reduction and adsorption) 1318-93-0 HCAPLUS RNMontmorillonite ((Al1.33-1.67Mg0.33-0.67)(Ca0-1Na0-1)0.33Si4(OH)2O10.xH2O) CN (9CI) (CA INDEX NAME) \*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\* 1344-28-1, Alumina, uses RL: CAT (Catalyst use); NUU (Other use, unclassified); USES (Uses) (hydrogenolysis catalyst support and adsorbent; purification of methylal containing peroxides by reduction and adsorption) 1344-28-1 HCAPLUS RN Aluminum oxide (Al2O3) (8CI, 9CI) (CA INDEX NAME) CN\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\* 7440-02-0, Raney nickel, uses 7440-05-3, Palladium, uses 7440-06-4, Platinum, uses RL: CAT (Catalyst use); USES (Uses) (hydrogenolysis catalyst; purification of methylal containing peroxides by reduction and adsorption) ΡN 7440-02-0 HCAPLUS CN Nickel (8CI, 9CI) (CA INDEX NAME) Νi 7440-05-3 HCAPLUS RNPalladium (8CI, 9CI) (CA INDEX NAME) CN

Pd

7440-06-4 HCAPLUS RNPlatinum (8CI, 9CI) (CA INDEX NAME) CN

Pt

CN

16940-66-2, Sodium borohydride IT 16971-29-2D, Borohydride, trimethoxy-, salts 33195-00-5D , Cyanoborohydride, salts RL: RGT (Reagent); RACT (Reactant or reagent) (reducing agent; purification of methylal containing peroxides by reduction and adsorption) 16940-66-2 HCAPLUS RNBorate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

• Na+

16971-29-2 HCAPLUS RNBorate(1-), tetrahydro- (8CI, 9CI) (CA INDEX NAME)

RN 33195-00-5 HCAPLUS CN Borate(1-), (cyano-κC)trihydro-, (T-4)- (9CI) (CA INDEX NAME)

L33 ANSWER 7 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2003:22609 HCAPLUS DOCUMENT NUMBER: 138:94840 Extraction and recovery of ions from a solution TITLE: INVENTOR(S): Strauss, Steven H.; Odom, Matthew A.; Clapsaddle, Brady PATENT ASSIGNEE(S):

Colorado State University Research Foundation, USA

SOURCE: PCT Int. Appl., 32 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT	NO.	K	ND	DATE		i	APPL	ICAT	ION 1	NO.		D	ATE	
		1				1	WO 2	002-	US20:	973		2	0020	627
	001895 AE, AG,			2003 , AU,		BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
	CO, CR,													
	GM, HR,													
	LS, LT,	LU, LV	, MA	, MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NO,	NZ,	OM,	PH,
	PL, PT,	RO, RU	J, SD	, SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TN,	TR,	TT,	TZ,
	UA, UG,	US, UZ	, VN	, YU,	ZA,	ZM,	ZW,	AM,	ΑZ,	BY,	KG,	KZ,	MD,	RU,
	TJ, TM													
RW:	GH, GM,	KE, LS	, MW	, MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AT,	BE,	CH,
	CY, DE,	DK, ES	FI,	, FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,
	BF, BJ,	CF, CC	, CI	, CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG
US 2005	145571	I	1	2005	0707	1	US 2	003-	5018	84		2	0020	627
PRIORITY API	LN. INFO	.:				1	US 2	001-	3020	52P	]	P 2	0010	629
						1	WO 2	002-	US20:	973	Ţ	W 2	0020	627

AB The present invention provides organometallic redox-recyclable solid ion-extractant and a method for using the same in extracting and recovering perfluoroalkyl carboxylate or perfluoroalkyl sulfonate ions in a solution The solid ion extractant is a composition of the formula [ZX1Si(X2R2)(X3R3)R1Ar1MAr2]Ya, wherein: each of Ar1 and Ar2 is independently C4-C20 aryl selected from the group consisting of cyclopentadienyl, dicarbollide and Ph, each of which can be optionally substituted; M is a transition metal is selected from Fe, Ru, Mn, Co, Ni, Cr, Os, Rh and Ir; R1 is C2-C20 alkylene; each of X1, X2 and X3 is independently a bond, O, S, or NR4; each of R2, R3 and R4 is independently H, or C1-C6 alkyl; Z is a solid support selected from glass, metal or a polymeric resin; Y is an anion selected from NO3-, halide, HSO4-, ClO4-, ReO4-, PF6-, carboxylate and CF3SO3-; and a is 0 when said ion coordinating agent is deactivated, and a is an integer from 1-3 when said ion coordinating agent is activated. In particular, the present invention provides a method for extracting and recovering ions which are relatively insol. in a solvent at a particular solvent temperature range. More specifically, the present invention provides a method for extracting and recovering perfluoroalkylsulfonates from an aqueous solution, such as wastewaters

from firefighting.

7440-06-4D, Platinum, silyl compds., complexes of IT

RL: CAT (Catalyst use); USES (Uses)

(hydrosilylation catalyst; preparation and use of a recyclable solid extractant for extraction and recovery of perfluoroalkyl carboxylate or perfluoroalkyl sulfonate ions from wastewater or other aqueous solns.)

7440-06-4 HCAPLUS RN

CN Platinum (8CI, 9CI) (CA INDEX NAME)

Pt

IT 16940-66-2, Sodium borohydride
 RL: CPS (Chemical process); PEP (Physical, engineering or chemical process); PROC (Process)
 (reductant to deactivate extractant; preparation and use of a recyclable solid extractant for extraction and recovery of perfluoroalkyl carboxylate or perfluoroalkyl sulfonate ions from wastewater or other aqueous solns.)
RN 16940-66-2 HCAPLUS
CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

● Na+

L33 ANSWER 8 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2002:827745 HCAPLUS

DOCUMENT NUMBER: 137:341465

TITLE: Method and apparatus for generation of clean air moist

for indoor air purification

INVENTOR(S): Suda, Seijiro

PATENT ASSIGNEE(S): Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2002317980	A2	20021031	JP 2001-117286	20010416
PRIORITY APPLN. INFO.:			JP 2001-117286	20010416
AB The method is carri	ed out	by contacti	ng alkaline agueous s	solution contai

AB The method is carried out by contacting alkaline aqueous solution containing metal-hydrogen complex compound (e.g., NaBH4) with H2-generation catalyst (e.g., fluorinated Ni-alloy) to form H2 gas, then mixing the H2 with O2-excess gas (e.g., air) in the presence of oxidation catalyst (e.g., Pd) for low-temperature combustion to produce air having water content, to be supplied to the indoor air via a pipeline.

IT 7440-05-3, Palladium, uses

RL: CAT (Catalyst use); DEV (Device component use); NUU (Other use, unclassified); USES (Uses)

(method and apparatus for generation of clean air moist for indoor air purification)

RN 7440-05-3 HCAPLUS

CN Palladium (8CI, 9CI) (CA INDEX NAME)

Pd

IT 16940-66-2, Sodium borohydride (NaBH4)

RL: NUU (Other use, unclassified); RCT (Reactant); RACT (Reactant or reagent); USES (Uses)

(method and apparatus for generation of clean air moist for indoor air purification)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

Na+

L33 ANSWER 9 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN 2002:638003 HCAPLUS ACCESSION NUMBER: 137:188572 DOCUMENT NUMBER: Acidic redox leaching for recovery of precious TITLE: metals from low-concentration sources INVENTOR(S): Farone, William A.; Azad, Maryam H. PATENT ASSIGNEE(S): USA U.S. Pat. Appl. Publ., 11 pp. SOURCE: CODEN: USXXCO DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE \_\_\_\_\_\_ A1 20020822 US 2001-789270 20010220 US 2002112569 20030422 US 6551378 B2 US 2001-789270 PRIORITY APPLN. INFO.: 20010220 The Au and Pt-group metals in powdered low-concentrate sources (especially ore tailings) are recovered by leaching with aqueous solution containing HCl and an oxidizing agent (typically Cl2), followed by separation of the leach solution, extraction of the associated base metals, and reduction with precipitation of the precious metals for recovery. The residual solids from acidic leaching are neutralized with a base, and most of the base metals are precipitated as hydroxides. The process is environmentally acceptable, uses no organic solvents, and is suitable for leaching of ore tailings and wastes containing ppm amts. of precious metals. 7440-06-4, Platinum, processes 7440-57-5, ΙT Gold, processes RL: EPR (Engineering process); PEP (Physical, engineering or chemical process); PROC (Process) (leaching of; acidic redox leaching for recovery of precious metals from low-concentration sources) 7440-06-4 HCAPLUS RN Platinum (8CI, 9CI) (CA INDEX NAME) CN Pt 7440-57-5 HCAPLUS RN CN Gold (8CI, 9CI) (CA INDEX NAME) Au

# IT 16940-66-2, Sodium borohydride

RL: CPS (Chemical process); PEP (Physical, engineering or chemical process); PROC (Process)

(reduction with, in leach solution; acidic redox leaching for recovery of precious metals from low-concentration sources)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

• Na+

L33 ANSWER 10 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:91311 HCAPLUS

DOCUMENT NUMBER: 134:118619

TITLE: Method for recovery of palladium from spent catalyst solution for electroless coating

INVENTOR(S): Nakamura, Tomonobu; Tsuchida, Kazunori

PATENT ASSIGNEE(S): Matsuda Sangyo K. K., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 7 pp. CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2001032025	A2	20010206	JP 1999-204878	19990719
PRIORITY APPLN. INFO.:			JP 1999-204878	19990719

AB The method comprises reducing with Na borohydride.

IT 7440-05-3P, Palladium, preparation

RL: PUR (Purification or recovery); PREP (Preparation) (recovery of palladium from spent catalyst solution for electroless coating by reducing with sodium borohydride)

RN 7440-05-3 HCAPLUS

CN Palladium (8CI, 9CI) (CA INDEX NAME)

Pd

IT 16940-66-2, Sodium borohydride

RL: RCT (Reactant); RACT (Reactant or reagent)
 (recovery of palladium from spent catalyst solution
 for electroless coating by reducing with sodium
 borohydride)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

● Na +

L33 ANSWER 11 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1998:42117 HCAPLUS

DOCUMENT NUMBER:

128:104629

TITLE:

Method for recovery of palladium from spent

catalysts

INVENTOR(S):

Nakazawa, Hiroyuki

PATENT ASSIGNEE(S):

Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 3 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 10008155	A2	19980113	JP 1996-192651	19960619
PRIORITY APPLN. INFO.:			JP 1996-192651	19960619

AΒ The method comprises dissolving a spent catalyst with aqua regia, precipitating the dissolved Pd as iodide for separation, and reducing the iodide with a reducing agent selected from NaBH4, hydrazine hydrochloride, hydrazine sulfate, hydrazine, Na2SO3, Na2S2O6, formic acid, oxalic acid.

IT 7440-05-3P, Palladium, preparation

RL: PUR (Purification or recovery); PREP (Preparation)

(recovery of palladium from spent catalysts by separation as

iodide and reduction)

7440-05-3 HCAPLUS RN

Palladium (8CI, 9CI) (CA INDEX NAME) CN

Pd

16940-66-2, Sodium borohydride IT

RL: TEM (Technical or engineered material use); USES (Uses) (recovery of palladium from spent catalysts by

separation as iodide and reduction)

RN 16940-66-2 HCAPLUS

Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME) CN

Na +

L33 ANSWER 12 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1997:93860 HCAPLUS

DOCUMENT NUMBER: 126:106889

TITLE: Separation and recovery of platinum-group metals in solutions with high efficiency

INVENTOR(S): Kawasaki, Hajime; Amino, Maki; Nishimura, Kenji

PATENT ASSIGNEE(S): Mitsubishi Materials Corp, Japan SOURCE: Jpn. Kokai Tokkyo Koho, 10 pp. CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 08291345	A2	19961105	JP 1995-95134	19950420
JP 3348808	B2	20021120		
ORITY APPLN. INFO.:			JP 1995-95134	19950420

PRIORITY APPLN. INFO.:

AB The process comprises (1) formation of complexes by adding complexing agents selected from ≥1 phosphines, phosphites, dialkylsulfides, cyclopentadiene, and/or CO in aqueous solns. or organic solns. containing Pt -group metal salts, (2) optional solid-liquid separation, (3) extraction of the complexes by contacting with supercrit. fluids, and (4) blowing into NH4OH aqueous solns. containing N2H4, NaBH4, thiourea, or NH4Cl for recovery of Pt-group metals. The phosphines may be selected from trimethylphosphine, triethylphosphine, tripropylphosphine, or tributylphosphine. The phosphites may be selected from trimethylphosphite, triethylphosphite, tripropylphosphite, or tributylphosphite. The dialkylsulfides may be selected from butylsulfide, hexylsulfide, or octylsulfide. The supercrit. fluids. may be selected from CO2, fluorocarbons, or N2O.

IT 16940-66-2, Sodium borohydride (NaBH4)

RL: NUU (Other use, unclassified); USES (Uses)
(blowing of ammonia aqueous solns. containing; platinum-group

metals separation and recovery by complexation and extraction
with supercrit. fluids)

with supercrit. Iluius

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

#### • Na+

NAME)

IT 7440-05-3P, Palladium, preparation

RL: PUR (Purification or recovery); PREP (Preparation) (platinum-group metals separation and recovery by complexation and extraction with supercrit. fluids)

RN 7440-05-3 HCAPLUS

CN Palladium (8CI, 9CI) (CA INDEX NAME)

Pd

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L33 ANSWER 13 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
                         1996:485373 HCAPLUS
DOCUMENT NUMBER:
                         125:201342
TITLE:
                         Precipitation behavior of metallic palladium
                         and platinum in aqueous solutions
AUTHOR (S):
                         Sakurai, Hiroki; Hirokawa, Kichinosuke
CORPORATE SOURCE:
                         Inst. Mater. Res., Tohoku Univ., Sendai, 980-77, Japan
                         Bunseki Kagaku (1996), 45(8), 795-798
SOURCE:
                         CODEN: BNSKAK; ISSN: 0525-1931
PUBLISHER:
                         Nippon Bunseki Kagakkai
DOCUMENT TYPE:
                         Journal
                         Japanese
LANGUAGE:
    Pd and Pt were reduced and precipitated in aqueous acidic solns.
    by sodium tetrahydroborate, sodium phosphinate and
     ascorbic acid. Pd was reduced by all the reducing agents, but
     Pt was reduced by sodium tetrahydroborate
     only. Both metal could be reduced individually, but when they
     were co-existed, complete separation could not be achieved by these reduction
for
     the generation of active hydrogen during reduction of Pd. The
     technic is applicable to recycle and recover Pt and
    Pd from exhaust gas-purification waste catalysts.
    7440-05-3P, Palladium, preparation 7440-06-4P,
    Platinum, preparation
    RL: CAT (Catalyst use); PEP (Physical, engineering or chemical process);
     PUR (Purification or recovery); PREP (Preparation); PROC (Process); USES
        (precipitation behavior of metallic palladium and platinum
        in aqueous acidic solns. for separation recovery)
     7440-05-3 HCAPLUS
ВN
     Palladium (8CI, 9CI) (CA INDEX NAME)
CN
Pd
RN
     7440-06-4 HCAPLUS
     Platinum (8CI, 9CI) (CA INDEX NAME)
CN
Pt
IT
     16940-66-2, Sodium tetrahydroborate
     RL: NUU (Other use, unclassified); USES (Uses)
        (reducing agent; precipitation behavior of metallic palladium and
        platinum in aqueous acidic solns. for separation recovery)
     16940-66-2 HCAPLUS
RN
     Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)
CN
```

● Na+

L33 ANSWER 14 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:148121 HCAPLUS

DOCUMENT NUMBER: 124:261743

TITLE: Preparation of N-(long-chain acyl)amino acids and

their salts

INVENTOR(S): Kaneko, Yohei; Nishimoto, Yoshifumi

PATENT ASSIGNEE(S): Kao Corp, Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 07330699 A2 19951219 JP 1994-120924 19940602
PRIORITY APPLN. INFO:: JP 1994-120924 19940602

OTHER SOURCE(S): MARPAT 124:261743

AB The title salts, useful as surfactants and bacteriostats, are prepared by hydrolysis of R1CONR2R3CN [COR1 = C8-22 (un)saturated aliphatic acyl; R2 = H, C1-3 linear or branched alkyl; R3 = C1-5 linear or branched alkylene] in the presence of basic substances, followed by treatment of the resulting R1CONR2R3CO2M (M = cation) (I) with metal hydrides, by treatment of the resulting I with oxidizing agents, or by hydrogenation of the resulting I in the presence of hydrogenation catalysts. The salts are further treated with mineral acids to adjust pH at 1-5 to give the corresponding acyl amino acids. These methods provide the products with no offensive odor and coloration. A mixture of 70 g N-lauroyl-β-aminopropionitrile, H2O, and an aqueous KOH solution was stirred at 90° for 15 h and the crude K N-lauroyl-β-alaninate (II) obtained was further treated with NaBH4 at 60° for 2 h to give 250 g aqueous solution containing 32.8% II with APHA color 100. After removal of NH3 from the solution

by evaporation, no offensive odor was detected.

IT 7440-02-0, Nickel, uses

RL: CAT (Catalyst use); USES (Uses)

(Raney; preparation of (higher acyl)amino acids (salts) by hydrolysis of amidonitriles followed by chemical purification)

RN 7440-02-0 HCAPLUS

CN Nickel (8CI, 9CI) (CA INDEX NAME)

Ni

IT 7440-05-3, Palladium, uses 7440-06-4,

Platinum, uses

RL: CAT (Catalyst use); USES (Uses)

(preparation of (higher acyl)amino acids (salts) by hydrolysis of amidonitriles followed by chemical purification)

RN 7440-05-3 HCAPLUS

CN Palladium (8CI, 9CI) (CA INDEX NAME)

Pd

RN 7440-06-4 HCAPLUS

CN Platinum (8CI, 9CI) (CA INDEX NAME)

Ρt

IT 11138-47-9, Sodium perborate 16940-66-2, Sodium
borohydride
RL: RCT (Reactant); RACT (Reactant or reagent)

RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation of (higher acyl)amino acids (salts) by hydrolysis of
amidonitriles followed by chemical purification)

RN 11138-47-9 HCAPLUS

CN Perboric acid, sodium salt (8CI, 9CI) (CA INDEX NAME)

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

• Na+

L33 ANSWER 15 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:964911 HCAPLUS

DOCUMENT NUMBER: 124:96495

TITLE: Process for removing oils and greases from industrial

wastewaters

Guess, Robert G. INVENTOR (S):

Romar Technologies, Inc., USA PATENT ASSIGNEE(S):

U.S., 12 pp. Cont.-in-part of U.S. Ser. No. 160,240, SOURCE:

abandoned. CODEN: USXXAM

DOCUMENT TYPE:

Patent.

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5462670	Α	19951031	US 1994-218891	19940328
US 5122279	Α	19920616	US 1991-682129	19910408
US 5298168	Α	19940329	US 1992-893978	19920603
US 5545331	Α	19960813	US 1995-386700	19950210
PRIORITY APPLN. INFO.:			US 1991-682129 A	2 19910408
			US 1992-893978 A	3 19920603
			US 1993-160240 B	2 19931202
			US 1994-218891 A	2 19940328

AB A process for removing dissolved oils and greases from an aqueous solution which

also may contain dissolved heavy metals is provided wherein the aqueous solution is mixed with a source of ferrous ion and dithionite ion in a first step at acidic pH to reduce and permit removal of solid heavy metal, is present and to sep. oils and greases from the aqueous solution The solution from the first step is reacted in a second step with hydroxide slurries obtained from third and fourth steps. A second step solution from the second step is reacted in a third step with an alkali composition and a third solution Optionally, the third solution is reacted with a chelating agent

for iron and an oxidizer in a fourth step. A solution of chelated iron from the fourth step, when practical is disposed of. Oils and greases are recovered from the first step such as by skimming, and the heavy metals are recovered. The method provides effective treatment of contaminated groundwater, landfill leachate, sewage, mine drainage, slag pile drainage, mine tailings drainage, plating wastes, and etching wastes.

7429-90-5, Aluminum, uses 7439-89-6, Iron, uses

7440-66-6, Zinc, uses 16940-66-2, Sodium

borohydride

RL: NUU (Other use, unclassified); USES (Uses) (recovering heavy metals, oils and greases from industrial wastewaters in multistage process)

7429-90-5 HCAPLUS RN

Aluminum (8CI, 9CI) (CA INDEX NAME) CN

Al

7439-89-6 HCAPLUS ВM

Iron (7CI, 8CI, 9CI) (CA INDEX NAME) CN

Fe

RN 7440-66-6 HCAPLUS CNZinc (7CI, 8CI, 9CI) (CA INDEX NAME)

Zn

16940-66-2 HCAPLUS RNBorate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME) CN

• Na+

IT7439-92-1P, Lead, preparation 7440-02-0P, Nickel, preparation 7440-05-3P, Palladium, preparation 7440-06-4P, Platinum, preparation 7440-22-4P, Silver, preparation 7440-31-5P, Tin, preparation 7440-48-4P, Cobalt, preparation 7440-50-8P, Copper, preparation 7440-57-5P, Gold, preparation RL: PUR (Purification or recovery); PREP (Preparation) (recovering heavy metals, oils and greases from industrial wastewaters in multistage process)

RN

7439-92-1 HCAPLUS Lead (8CI, 9CI) (CA INDEX NAME) CN

Pb

RN7440-02-0 HCAPLUS CN Nickel (8CI, 9CI) (CA INDEX NAME)

Νi

7440-05-3 HCAPLUS RNPalladium (8CI, 9CI) (CA INDEX NAME) CN

Pđ

RN 7440-06-4 HCAPLUS CN Platinum (8CI, 9CI) (CA INDEX NAME)

Pt

7440-22-4 HCAPLUS RN

Silver (8CI, 9CI) (CA INDEX NAME) CN

Ag

RN7440-31-5 HCAPLUS

CNTin (8CI, 9CI) (CA INDEX NAME)

Sn

RN7440-48-4 HCAPLUS

Cobalt (8CI, 9CI) (CA INDEX NAME) CN

Co

RN7440-50-8 HCAPLUS

Copper (7CI, 8CI, 9CI) (CA INDEX NAME) CN

Cu

RN

7440-57-5 HCAPLUS Gold (8CI, 9CI) (CA INDEX NAME) CN

Au

L33 ANSWER 16 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1995:563523 HCAPLUS
DOCUMENT NUMBER: 122:295785
TITLE: Ore leaching with non-cyanide redox solution for recovery of precious metals
INVENTOR(S): Martenson, Irvin; Mathis, Samuel A.
PATENT ASSIGNEE(S): USA

SOURCE: U.S., 6 pp.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5401296	Α	19950328	US 1994-266563	19940628
CA 2194137	AA	19960111	CA 1995-2194137	19950324
WO 9600801	A1	19960111	WO 1995-US3734	19950324
W: AU, BR, CA,	CN, MX	, RU		
AU 9521949	A1	19960125	AU 1995-21949	19950324
ZA 9502471	A	19960415	ZA 1995-2471	19950327
PRIORITY APPLN. INFO.:			US 1994-266563 A	19940628
			WO 1995-US3734 W	19950324

WO 1995-US3734 W 19950324

AB The aqueous slurry with 20-25% ore powder (size -250 mesh) for leaching of Au, Ag, and/or Pt-group metals is based on the aqueous solution containing NaBr 1.5-3.5, NaCl 3-6%, K2S208 for the redox potential of 700-900 mV, and HCl for the pH of 3-4. The resulting leach liquor is pumped to a holding tank, and reacted with aqueous solution of NaOH and NaBH4 at pH of 8.2-8.6 to precipitate the precious metals for recovery by filtration, followed by conventional separation and purification The liquor from filtration is

electrochem. treated with replenishment of the salts, and recycled to the leaching stage. The K2S2O8 is preferably com. Oxone triple salt. The redox leaching process is suitable for recovery of Pt, Ir, and Pd from volcanic ash.

IT 7440-05-3P, Palladium, preparation 7440-06-4P,

Platinum, preparation 7440-22-4P, Silver, preparation

7440-57-5P, Gold, preparation

RL: PUR (Purification or recovery); PREP (Preparation)

(leaching of; ore leaching with non-cyanide redox solution for recovery of precious metals)

RN 7440-05-3 HCAPLUS

CN Palladium (8CI, 9CI) (CA INDEX NAME)

Pd

RN 7440-06-4 HCAPLUS

CN Platinum (8CI, 9CI) (CA INDEX NAME)

Ρt

RN 7440-22-4 HCAPLUS

CN Silver (8CI, 9CI) (CA INDEX NAME)

Ag

7440-57-5 HCAPLUS RNGold (8CI, 9CI) (CA INDEX NAME) CN

Au

16940-66-2, Sodium borohydride (NaBH4) IT RL: MOA (Modifier or additive use); USES (Uses) (redox solution containing; ore leaching with non-cyanide redox solution for recovery of precious metals)

16940-66-2 HCAPLUS RN

Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME) CN

• Na+

L33 ANSWER 17 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1994:196570 HCAPLUS DOCUMENT NUMBER: 120:196570 TITLE: Rhodium, platinum, and palladium recovery from new and spent automotive catalysts AUTHOR (S): Wu, Kuo Ying Amanda; Wisecarver, Keith D.; Abraham, Martin A.; Takach, Nicholas; Yang, Ningsheng Dep. Chem. Eng., Univ. Tulsa, Tulsa, OK, 74104, USA CORPORATE SOURCE: Precious Metals (1993), 17th, 343-9 SOURCE: CODEN: PRCMEU; ISSN: 8756-0917 DOCUMENT TYPE: Journal LANGUAGE: English The recovery of Pt, Pa, and Rh automotive catalysts by leaching HCl + H2SO4 with the addition of F- and H2O2 including pretreatment with Na borohydride, sodium formate and NaOH was studied. The recovery of metals mainly depends on the catalyst type and occurs in the following sequence Pd>Pt >Rh. Rh recovery of 90% or better can be achieved from both new and spent catalysts using pretreatment with sodium formate or NaOH. 16940-66-2, Sodium borohydrate TT RL: PROC (Process) (recovery of palladium and platinum and rhodium from spent automotive catalyst by leaching with acids and pretreatment with) 16940-66-2 HCAPLUS RN

Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

CN

## Na +

RN 7440-06-4 HCAPLUS

CN Platinum (8CI, 9CI) (CA INDEX NAME)

Ρt

L33 ANSWER 18 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1994:57151 HCAPLUS DOCUMENT NUMBER: 120:57151 Recovery of noble metals from solutions TITLE: produced in oxidation reactions involving alkyl nitrites by treatment with reducing agents Landscheidt, Heinz; Klausener, Alexander; Blank, Heinz INVENTOR (S): Ulrich Bayer A.-G., Germany PATENT ASSIGNEE(S): Ger. Offen., 6 pp. SOURCE: CODEN: GWXXBX DOCUMENT TYPE: Patent German LANGUAGE: FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. -------------------A1 19930527 DE 1991-4137965 DE 4137965 19911119 A1 19930616 B1 19950419 19930616 EP 1992-119030 EP 546324 19921106 EP 546324 R: AT, BE, CH, DE, FR, GB, IT, LI, NL AT 121316 E 19950515 AT 1992-119030
JP 05239567 A2 19930917 TO 2007 19921106 19921117 DE 1991-4137965 A 19911119 PRIORITY APPLN. INFO.: Wastewaters containing metals or metal salts are treated with a reducing agent and an additive to reduce the metals, which are recovered in the metallic form. Suitable reducing agents include H2, CO, or a complex hydride, preferably a boron hydride of Group I or II metals, especially Na, K, Li, or Zn. The additive is a primary or secondary amino group which can react with nitrous acid, e.g., ammonia, an ammonium salt, ammonium chloride, Me ammonium chloride, urea, or amidosulfonic acid. The metallic form can be changed to a salt form by oxidation with an oxidizing agent in an aqueous acid solution, e.g., HCl or acid. Suitable oxidizing agents include chlorates or nitrates of alkali or alkaline earth metals or ammonium ion, HNO3, or H2O2. The method is suitable for recovering Pt group metals, especially IT 7647-10-1, Palladium chloride RL: CAT (Catalyst use); USES (Uses) (catalysts, for Me acrylate reaction with Me nitrite, recovery of) RN7647-10-1 HCAPLUS CN Palladium chloride (PdCl2) (6CI, 8CI, 9CI) (CA INDEX NAME) C1-Pd-C1 IT 7440-05-3P, Palladium, preparation RL: PREP (Preparation) (recovery of, from oxidation reaction mixts.)

RN7440-05-3 HCAPLUS CN Palladium (8CI, 9CI) (CA INDEX NAME)

Pđ

13762-51-1 16940-66-2 16949-15-8, IT

Lithium boron hydride 17611-70-0

RL: USES (Uses)

(reducing agents, in recovery of noble metal
catalysts)

RN 13762-51-1 HCAPLUS

CN Borate(1-), tetrahydro-, potassium (8CI, 9CI) (CA INDEX NAME)

• K+

RN 16940-66-2 HCAPLUS CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

• Na+

RN 16949-15-8 HCAPLUS CN Borate(1-), tetrahydro-, lithium (8CI, 9CI) (CA INDEX NAME)

● Li+

RN 17611-70-0 HCAPLUS CN Borate(1-), tetrahydro-, zinc (2:1) (9CI) (CA INDEX NAME)

●1/2 Zn<sup>2+</sup>

L33 ANSWER 19 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN 1994:12294 HCAPLUS ACCESSION NUMBER: 120:12294 DOCUMENT NUMBER: Recovery of platinum-group metals TITLE: from metal foil support catalysts Ito, Hidetoshi; Eto, Yoshuki INVENTOR(S): PATENT ASSIGNEE(S): Nissan Motor, Japan SOURCE: Jpn. Kokai Tokkyo Koho, 3 pp. CODEN: JKXXAF DOCUMENT TYPE: Patent Japanese LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: DATE APPLICATION NO. DATE PATENT NO. KIND --------------JP 05212296 A2 19930824 JP 1992-17614 19920203 PRIORITY APPLN. INFO.: JP 1992-17614 19920203 The recovery involves (1) soaking the metal foil-supported catalysts in solns. containing NaOH ≥20, NaHSO4 1-10, and Na boron hydride ≥0.1 weight%, (2) dissolving the Al2O3- and CeO2 coating layers (as catalyst active layer), and (3) recovering the Pt -group metals as platinum black. The method can recover the metals efficiently, and the method is applied to catalysts for such as exhaust gas purification 1344-28-1, Alumina, uses IT RL: USES (Uses) (coat layers, catalysts containing, metal foil-support, recovery of platinum-group metals from) 1344-28-1 HCAPLUS RN Aluminum oxide (Al2O3) (8CI, 9CI) (CA INDEX NAME) \*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\* TΨ 16940-66-2 RL: PROC (Process) (dissoln. of alumina and ceria coat layers in solns. containing, in recovery recovery of platinum-group metals from metal foil-support catalysts) 16940-66-2 HCAPLUS RNBorate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME) CN 3+ Na+

CN Palladium (8CI, 9CI) (CA INDEX NAME)

Pd

RN 7440-06-4 HCAPLUS

CN Platinum (8CI, 9CI) (CA INDEX NAME)

Pt

L33 ANSWER 20 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN 1992:637544 HCAPLUS ACCESSION NUMBER: 117:237544 DOCUMENT NUMBER: Precious metal leaching with iodine-iodide TITLE: solutions Sloan, Hilbert INVENTOR(S): PATENT ASSIGNEE(S): Shapiro, Nelson H., USA U.S., 11 pp. Cont. of U.S. Ser. No. 69,100, abandoned. SOURCE: CODEN: USXXAM Patent DOCUMENT TYPE: English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: KIND DATE APPLICATION NO. DATE PATENT NO. \_ \_ \_ \_ US 5137700 US 1989-342587 19890424 Α 19920811 PRIORITY APPLN. INFO.: US 1987-69100 B1 19870702 Precious metals (especially Au, Ag, Pd, and Pt) are recovered from electroplated scrap or articles by immersion into aqueous etching bath containing I 1, NH4I (or HI) 4, and water 10 parts. The precious metals are oxidized by I, form a soluble complex with the iodide, and can be precipitated from the solution by addition of a weak reducing agent (especially hydroxylamine or NaBH4) and recovered by filtration. The filtrate is oxidized with H2O2 or (NH4)S2O8 to precipitate elemental I for recycling to the etching stage. 7440-05-3P, Palladium, preparation 7440-06-4P, TT Platinum, preparation 7440-22-4P, Silver, preparation 7440-57-5P, Gold, preparation RL: PUR (Purification or recovery); PREP (Preparation) (recovery of, from electroplated scrap, leaching with iodine-iodide solution for) 7440-05-3 HCAPLUS RN Palladium (8CI, 9CI) (CA INDEX NAME) CN Pd RN7440-06-4 HCAPLUS Platinum (8CI, 9CI) (CA INDEX NAME) CN Pt 7440-22-4 HCAPLUS RN Silver (8CI, 9CI) (CA INDEX NAME) CNAq RN 7440-57-5 HCAPLUS

Au

Gold (8CI, 9CI) (CA INDEX NAME)

IT 16940-66-2, Sodium borohydride
 RL: PROC (Process)
 (reducing agent, precious metal recovery with,
 after leaching with aqueous iodine-iodide solution)
RN 16940-66-2 HCAPLUS
CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

● Na+

L33 ANSWER 21 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1992:595687 HCAPLUS

DOCUMENT NUMBER:

117:195687

TITLE:

Recovery of palladium from

tetraamminepalladium dichloride solutions by reduction

Suzuki, Masahito; Nagai, Makoto

PATENT ASSIGNEE(S):

N. E. Chemcat Corp., Japan Jpn. Kokai Tokkyo Koho, 3 pp.

SOURCE:

CODEN: JKXXAF

DOCUMENT TYPE:

INVENTOR(S):

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND DATE ---------\_\_\_\_\_ -----JP 04165030 A2 19920610 JP 1990-287703 19901025

PRIORITY APPLN. INFO.:

JP 1990-287703

19901025

[Pd(NH3)4]Cl2 solns. are added to solns. containing a strong alkaline reducer. Agglomeration-free Pd powders are prepared without loss.

7440-05-3P, Palladium, preparation IT

RL: PREP (Preparation)

(recovery of powdered, from tetraamminepalladium chloride, by reduction with alkaline reducer)

7440-05-3 HCAPLUS RN

CN Palladium (8CI, 9CI) (CA INDEX NAME)

Pd

16940-66-2, Sodium borohydride IT

> RL: RCT (Reactant); RACT (Reactant or reagent) (reduction by, of tetraamminepalladium chloride, for recovery of powdered palladium)

16940-66-2 HCAPLUS RN

Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME) CN

#### Na +

13815-17-3 TT

RL: RCT (Reactant); RACT (Reactant or reagent) (reduction of, by alkaline reducer, for recovery of powdered palladium)

13815-17-3 HCAPLUS

Palladium(2+), tetraammine-, dichloride, (SP-4-1)- (9CI) (CA INDEX NAME) CN

$$^{\mathrm{NH_3}}_{|2+}$$
 $_{\mathrm{H_3N-Pd}}^{\mathrm{NH_3}}$ 
 $_{\mathrm{NH_3}}^{\mathrm{NH_3}}$ 

●2 C1-

L33 ANSWER 22 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:189660 HCAPLUS

DOCUMENT NUMBER: 114:189660

TITLE: Reducing agent for recovery of precious metals

from cyanide solutions

INVENTOR(S): Shoda, Toshiaki

PATENT ASSIGNEE(S): Nippon Electro Plating Engineers K. K., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 3 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

KIND DATE APPLICATION NO. DATE PATENT NO. \_ \_ \_ \_ \_\_\_\_\_ \_\_\_\_\_ \_\_\_\_\_ A2 19900622 JP 1988-316335 JP 02163327 19881216 JP 1988-316335 19881216 PRIORITY APPLN. INFO.: The loaded cyanide feed solution (especially from leaching) is reduced with powdered

Al, powdered Zn, N2H4, alkali metal salt of HB, H2PO3, and/or dithionous acid, and/or a carbonyl compound to recover the precious metal. Thus, the aqueous solution (pH 13) containing 50 g KCN/L and 4.3 g Au/L was mixed with powdered Al 0.2, HCHO 1.0, and N2H4 0.1M, and showed the residual Au of 2.5 after 1 day and and 1.6 mg/L after 5 days.

IT 7440-57-5P, Gold, preparation

RL: PUR (Purification or recovery); PREP (Preparation) (recovery of, from cyanide solution, reducing agents for)

RN 7440-57-5 HCAPLUS

CN Gold (8CI, 9CI) (CA INDEX NAME)

Au

IT 16940-66-2 7429-90-5, Aluminum, uses and miscellaneous 7440-66-6, Zinc, uses and miscellaneous

7440-00-0, Zinc, uses and miscerianeous

RL: PROC (Process)

(reducing agent, gold recovery with, from aqueous cyanide solution)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

• Na+

RN 7429-90-5 HCAPLUS

CN Aluminum (8CI, 9CI) (CA INDEX NAME)

Al

RN

7440-66-6 HCAPLUS Zinc (7CI, 8CI, 9CI) (CA INDEX NAME) CN

Zn

7440-50-8, Copper, uses and miscellaneous IT

RL: USES (Uses)

(reducing agent, gold recovery with, from cyanide solution) 7440-50-8 HCAPLUS

RN

Copper (7CI, 8CI, 9CI) (CA INDEX NAME) CN

Cu

L33 ANSWER 23 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:16564 HCAPLUS

DOCUMENT NUMBER: 114:16564

TITLE: Separation of rhodium(III) from solutions, containing

iridium(IV) and -(III)

AUTHOR(S): Volkov, A. A.; Khain, V. S.

CORPORATE SOURCE: Ukhtinsk. Ind. Inst., Ukhta, USSR

SOURCE: Zhurnal Neorganicheskoi Khimii (1990), 35(9), 2220-3

CODEN: ZNOKAQ; ISSN: 0044-457X

DOCUMENT TYPE: Journal LANGUAGE: Russian

AB A mixture of Ir(IV) as IrCl62- or Ir(III) as IrCl63- with Rh(III) as

Rh(OH)63- was separated in strong acidic media (pH < 1) by NaBH4. The method

can be used in refining Pt-group metals and for their

concentration with high yield and purity.

IT 16940-66-2, Sodium tetrahydroborate

RL: RCT (Reactant); RACT (Reactant or reagent)

(reduction by, of trivalent rhodium or tetravalent or trivalent iridium in

separation of rhodium from iridium)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

● Na+

L33 ANSWER 24 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1990:408958 HCAPLUS

DOCUMENT NUMBER: 113:8958

TITLE: Chemical separation for platinum from

rhodium

INVENTOR(S): Zhang, Jian; Xu, Ying; Zhang, Ji

PATENT ASSIGNEE(S): Xian Non-Ferrous Metals Institute, Peop. Rep. China SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, 5 pp.

CODEN: CNXXEV

DOCUMENT TYPE: Patent LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1034143	Α	19890726	CN 1988-100131	19880111
CN 1011599	В	19910213		

PRIORITY APPLN. INFO.: CN 1988-100131 19880111

AB In electrolysis of Pt-Rh alloy, a small amount of H2O2 is added to the electrolyte solution for increasing the dissolving of Pt-Rh alloy electrodes, and the resulting electrolyte solution is then mixed with KCl to form Pt compound precipitate The Pt compound precipitate is reduced with NaBH4 to recover ≥99% pure Pt. Thus, waste Pt-Rh alloy chips were pressed to form electrodes, immersed into 8N HCl, placed in an electrolysis cell with magnetic stirrer, added 6 mL H2O2, and then electrolyzed at 180 mA/cm2 for 3 h. The electrolyte solution containing 72.568 g/L Rn and 8.063 g/L Pt was mixed with 120% KCl, precipitated and filtered to obtain yellow precipitate, which was then washed with 10%

KCl and 5% HCl solns., dried, and reduced with NaBH4 to recover 99.5% pure **Pt.** The filtrate was passed through an ion-exchange resin bed to recover 99.9% pure Rh.

IT 16940-66-2

RL: USES (Uses)

(reducing agent, for platinum recovery, in electrolysis of platinum-rhodium alloy)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

#### Na +

IT 7440-06-4P, Platinum, preparation

RL: PREP (Preparation)

(separation of, from **platinum**-rhodium alloy, by electrolysis, hydrogen peroxide addition in)

RN 7440-06-4 HCAPLUS

CN Platinum (8CI, 9CI) (CA INDEX NAME)

Pt

L33 ANSWER 25 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1989:621064 HCAPLUS

DOCUMENT NUMBER: 111:221064

TITLE: Recovering of noble metal from solution INVENTOR(S): Yamazaki, Hiroshi; Tateno, Yoshihisa

PATENT ASSIGNEE(S): Tanaka Noble Metal Industrial Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 3 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01083682	A2	19890329	JP 1987-241755	19870926
PRIORITY APPLN. INFO.:			JP 1987-241755	19870926

AB A method for recovering a noble metal contained in a solution at least partially as a complex involves electrolysis to reduce the noble metal ions followed by recovery as well as to decompose the complex and recovering the noble metal ions formed by the decomposition using a reducing agent.

IT 7440-05-3P, Palladium, preparation 7440-06-4P,

Platinum, preparation
RL: PREP (Preparation)

(recovery of, by electrolysis and reducing agents)

RN 7440-05-3 HCAPLUS

CN Palladium (8CI, 9CI) (CA INDEX NAME)

Pd

RN 7440-06-4 HCAPLUS

CN Platinum (8CI, 9CI) (CA INDEX NAME)

Pt

IT 16940-66-2, Sodium borohydride

RL: PRP (Properties)

(reducing agents, in recovery of noble metals)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

Na +

L33 ANSWER 26 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1989:599088 HCAPLUS

DOCUMENT NUMBER:

111:199088

TITLE:

Recovery of platinum-group metals

from spent catalysts

INVENTOR(S):

Ezawa, Nobuyasu

PATENT ASSIGNEE(S):

Tanaka Noble Metal Industrial Co., Ltd., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

1

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01108323	A2	19890425	JP 1987-264961	19871020
PRIORITY APPLN. INFO.:			JP 1987-264961	19871020

Spent catalysts of supported type containing Pt-group metals AB and/or their oxides are treated to recover the metal values.

The process comprises: (1) reduction with NaBH4; (2) washing for Na removal; (3) chlorination leaching with aqueous HCl and sparged Cl; (4) leaching with aqueous HCl; (5) fluidized-bed treatment for cathodic recovery of Pt -group metals on C particles in an electrolysis cell; and (6)

recirculating the spent solns. for addnl. leaching.

16940-66-2, Sodium borohydride (NaBH4) IT

RL: PROC (Process)

(reduction with, of spent catalysts containing metal oxides in platinum-group metal recovery)

RN

16940-66-2 HCAPLUS
Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME) CN

Na +

L33 ANSWER 27 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1989:410611 HCAPLUS DOCUMENT NUMBER: 111:10611 TITLE: Recovery of platinum-group metal from spent electrode Sato, Hiroyasu; Maeda, Osamu INVENTOR(S): Tanaka Noble Metal Industrial Co., Ltd., Japan PATENT ASSIGNEE(S): SOURCE: Jpn. Kokai Tokkyo Koho, 3 pp. CODEN: JKXXAF DOCUMENT TYPE: Patent LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION: KIND DATE PATENT NO. APPLICATION NO. DATE \_\_\_\_ -----\_\_\_\_\_\_ JP 01008228 A2 19890112 JP 1987-163182 19870630 B2 19970129 JP 2575715 JP 1987-163182 PRIORITY APPLN. INFO.: 19870630 A spent electrode from NaCl electrolysis is treated to recover Pt -group metals by reduction of the PdO2, PtO2, and/or RhO2 layer on the surface with a reducing agent (especially HCHO) followed by leaching. Thus, a spent electrode from Ti mesh coated with 6:4 (mol) Pt-PdO2 layer was reduced with aqueous 0.5% N2H2.2HCl, and then leached with 6:1 HCl-HNO3 mixture at 40-80°. The leached electrode was recoated with Pt and PdO2, and then used in NaCl electrolysis. 7440-05-3P, Palladium, preparation 7440-06-4P, IT Platinum, preparation RL: PUR (Purification or recovery); PREP (Preparation) (recovery of, from spent electrode in salt electrolysis, reduction and leaching in) RN7440-05-3 HCAPLUS Palladium (8CI, 9CI) (CA INDEX NAME) CN Ρđ RN7440-06-4 HCAPLUS Platinum (8CI, 9CI) (CA INDEX NAME) CN Pt 16940-66-2 TT RL: PROC (Process) (reduction with, for platinum-group metal recovery from spent salt-electrolysis electrode) RN16940-66-2 HCAPLUS

Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

CN

● Na+

L33 ANSWER 28 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1989:158128 HCAPLUS

DOCUMENT NUMBER: 110:158128

TITLE: Recovery of platinum-group metals

from spent electrodes

INVENTOR(S): Sato, Hiroyasu; Maeda, Osamu

PATENT ASSIGNEE(S): Tanaka Noble Metal Industrial Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 3 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE \_\_\_\_\_ ----\_\_\_\_\_ \_\_\_\_\_\_ A2 19881108 JP 63270421 JP 1987-104133 19870427 JP 1987-104133 19870427 PRIORITY APPLN. INFO.:

AB Spent NaCl-electrolysis electrodes coated with a Pt-group metal oxide are treated to recover Pt, Ir, and Ru. The

oxide coating is reduced, leached with aqueous solution containing HC1 and HNO3 to

recover a part of Pt and Ir, and then electrolyzed in a fluoride bath for coating separation Suitable reducing agents are N2H4, NaBH4, and sulfites, and the fluoride is HF and/or NH4HF2. Thus, spent Ti electrodes had a coating with 88:12:5 mol mixture of Pt oxide, Ir2O3, and Ru oxide. The electrodes were dipped for 1 h in aqueous 0.2% N2H4·2HCl, and then for 1.5 h in 6:1 HCl:HNO3 mixture at 60-70°. The treated electrodes as anode were electrolyzed at 0.2 A/dm2 and 4.2 V for 10 min in aqueous bath containing 5 weight% each of HF and NH4HF2. The Pt, Ir, and Ru were recovered.

IT 7440-06-4P, Platinum, preparation

RL: PUR (Purification or recovery); PREP (Preparation)
(recovery of, from spent electrolysis electrodes, reduction of oxide coating for electrochem.)

RN 7440-06-4 HCAPLUS

CN Platinum (8CI, 9CI) (CA INDEX NAME)

Pt

IT 16940-66-2

RL: PROC (Process)

(reduction with, of metal oxide coating on spent electrode, recovery of platinum-group metals after)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

● Na+

L33 ANSWER 29 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1989:158127 HCAPLUS

DOCUMENT NUMBER: 110:158127

TITLE: Recovery of platinum-group metals

from spent electrodes

INVENTOR(S): Sato, Hiroyasu; Maeda, Osamu

PATENT ASSIGNEE(S): Tanaka Noble Metal Industrial Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 3 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 63270420 A2 19881108 JP 1987-104132 19870427

PRIORITY APPLN. INFO:: JP 1987-104132 19870427

AB Spent NaCl-electrolysis electrodes having a Pt-group metal oxide coating are treated to recover Pt, In, and

Ru. The oxide coating is reduced, leached with aqueous solution containing HCl and

HNO3 to recover a part of Pt and In, and then contacted with a mixture of HNO3, H2SO4, and HF for dissoln. Suitable reducing agents are N2H4, N2H4, NaBH4, and sulfites. Thus, spent Ti electrodes had a coating with 88:12:5 mol mixture of Pt oxide, In2O3, and Ru oxide. The electrodes were treated for 1 h in aqueous 0.2% N2H4.2HCl, for 1.5 h in 6:1 HCl:HNO3 solution at 60-70°, and then for 10 min at 50-60° in a solution containing HNO3 2, H2SO4 2, HF 1, and water 5 volume parts to recover Pt, Ir, and Ru.

IT 7440-06-4P, Platinum, preparation

RL: PUR (Purification or recovery); PREP (Preparation)
(recovery of, from spent electrodes, reduction of oxide coatings for leaching in)

RN 7440-06-4 HCAPLUS

CN Platinum (8CI, 9CI) (CA INDEX NAME)

Pt

IT 16940-66-2

RL: PROC (Process)

(reduction with, of oxide coating on spent electrode, recovery of platinum-group metals after)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

• Na+

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L33 ANSWER 30 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN
                        1986:37545 HCAPLUS
ACCESSION NUMBER:
                        104:37545
DOCUMENT NUMBER:
                        Winning precious metals from geological
TITLE:
                        brines
                        Luebbe, Ray H., Jr.; Wemhoff, Mark
INVENTOR(S):
                        Environmental Technology (U.S.), Inc., USA
PATENT ASSIGNEE(S):
                         Eur. Pat. Appl., 22 pp.
SOURCE:
                         CODEN: EPXXDW
                         Patent
DOCUMENT TYPE:
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
                                                                  DATE
                        KIND
                               DATE
                                           APPLICATION NO.
     PATENT NO.
                                           ______
                         ----
                                -----
                                                                  19850404
                                           EP 1985-302382
                         A2
                                19851030
     EP 159842
        R: DE, FR, GB
                                            US 1984-601792
                                                               A 19840419
PRIORITY APPLN. INFO.:
                                                               A 19840515
                                           US 1984-610385
     Geothermal and oil-field brines are contacted with an inorg. glassy
ΔR
     material for selective adsorption of precious metals, especially after
     surface pretreatment by reduction The metal-loaded glass is leached
     (e.g. with aqua regia) and recycled. Hydrophobic organic plastics and clays
     (especially attapulgite) are used as adsorbents. The brines are optionally
     pretreated with an oxidizing agent to enhance recovery of precious
     metals at less than +200 mV (redox) to avoid sorption of S and
     base elements. Thus, oil-field brine 5 gal was circulated at 2.4 L/min
     for 16 h through a glass chromatog. column packed with glass beads of 4 mm
     diameter The beads were washed with deionized water 125 mL of pH 9.5, and 5
     mL of cyanogen iodide solution was added. Recovery of Ag was only 0.03 mg/L,
     but was increased to 0.3 mg/L by extraction with aqua regia in 1:1 solution
     1309-48-4, uses and miscellaneous 1344-28-1, uses and
IT
     miscellaneous
     RL: USES (Uses)
        (activated, for adsorption of precious metals from brines)
     1309-48-4 HCAPLUS
RN
     Magnesium oxide (MgO) (9CI) (CA INDEX NAME)
CN
Mg = 0
     1344-28-1 HCAPLUS
RN
     Aluminum oxide (Al2O3) (8CI, 9CI) (CA INDEX NAME)
CN
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     7440-05-3P, preparation 7440-06-4P, preparation
IT
     7440-22-4P, preparation 7440-57-5P, preparation
     RL: PUR (Purification or recovery); PREP (Preparation)
        (recovery of, from brines, glassy sorbents for)
     7440-05-3 HCAPLUS
RN
     Palladium (8CI, 9CI) (CA INDEX NAME)
CN
Pd
```

RN

CN

7440-06-4 HCAPLUS

Platinum (8CI, 9CI) (CA INDEX NAME)

recovery from brines)
16971-29-2 HCAPLUS
Borate(1-), tetrahydro- (8CI, 9CI) (CA INDEX NAME)

RN CN

L33 ANSWER 31 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

1985:508225 HCAPLUS ACCESSION NUMBER:

103:108225 DOCUMENT NUMBER:

TITLE: Recovery of platinum metals

especially ruthenium

PATENT ASSIGNEE(S): Daido Steel Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 3 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent Japanese LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

F	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
-					
J	JP 60092433	A2	19850524	JP 1983-199154	19831026
	TY APPLN. INFO.:			JP 1983-199154	19831026
AB S	Spent electrodes or	cataly	sts are fuse	d with KOH + KNO3 (or	K2S2O7),
S	stirred in water, a	nd filt	ered. The f	iltrate is adjusted to	pH 5-6, and
f	filtered. The preci	ipitate	is dissolve	d in dilute HCl. filte	red, and the

filtered. The precipitate is dissolved in dilute HCl, filtered, and th filtrate

reduced with NaBH4 to obtain the Pt-group metals.

Thus, spent electrodes 15 g were processed with KOH + KNO3, to recover 23 mg metal containing Ru 87 and Ir 3.8%. The residue was heated at 800° to recover 56 mg sublimate containing 70% RuO2.

IT16940-66-2

RL: PROC (Process)

(reduction with, in processing of spent catalysts and electrodes for platinum-group metal recovery)

16940-66-2 HCAPLUS RN

Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME) CN

Na +

L33 ANSWER 32 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1983:581262 HCAPLUS

DOCUMENT NUMBER: 99:181262

TITLE:

Oxygen scavenging

PATENT ASSIGNEE(S): SOURCE: Kurita Kogyo Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 3 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 58079590	A2	19830513	JP 1981-177765	19811105
TD 50000105		1004000		

JP 59032195 B4 19840807

PRIORITY APPLN. INFO.: JP 1981-177765 19811105

AB An O-containing water is treated with a Pd catalyst in the presence of NaBH4 and/or N2H4 (or its hydrate). The Pd catalyst is loaded on an anion exchanger. Thus, a Pd chloride was treated with Lewatit OC 1045 [78769-27-4] to prepare a Pd catalyst. Boiler water was mixed with NaBH4 and then passed through a column containing the Pd-anion exchanger. The dissolved O in the eluent was 0.1 mg/L after 200 L water/h-L resin.

IT 7440-05-3, uses and miscellaneous

RL: CAT (Catalyst use); USES (Uses)

(catalyst, for oxygen scavenging from boiler water, Lewatit OC 1045 support for)

RN 7440-05-3 HCAPLUS

CN Palladium (8CI, 9CI) (CA INDEX NAME)

Pd

IT 16940-66-2

RL: OCCU (Occurrence)

(in oxygen scavenging in boiler water)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

Na +

L33 ANSWER 33 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN 1983:164610 HCAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 98:164610 The treatment of spent electroless nickel baths by TITLE: autocatalytic reduction AUTHOR(S): Parker, Konrad CORPORATE SOURCE: Park Ridge, IL, 60068, USA SOURCE: Plating and Surface Finishing (1983), 70(3), 60-2 CODEN: PSFMDH; ISSN: 0360-3164 DOCUMENT TYPE: Journal English LANGUAGE: The autocatalytic decomposition of some spent electroless Ni baths using NaBH4 and/or Pd ions can reduce the Ni concentration to  $\leq 10$  mg/L. The chemical of the plating solution appears to be a controlling factor. IT7440-02-0P, preparation RL: PUR (Purification or recovery); PREP (Preparation) (recovery of, from spent electroless bath, by autocatalytic reduction) RN7440-02-0 HCAPLUS Nickel (8CI, 9CI) (CA INDEX NAME) Νi IT 7647-10-1 16940-66-2 RL: RCT (Reactant); RACT (Reactant or reagent) (reduction by, of spent electroless bath, for nickel recovery) RN7647-10-1 HCAPLUS Palladium chloride (PdCl2) (6CI, 8CI, 9CI) (CA INDEX NAME) CNCl-Pd-Cl 16940-66-2 HCAPLUS RN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME) CN

● Na+

3+

L33 ANSWER 34 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1982:620285 HCAPLUS

DOCUMENT NUMBER: 97:220285

TITLE: Applications for sodium borohydride

in precious metal recovery and

recycle

AUTHOR(S): Medding, G. L.; Lander, J. A.

CORPORATE SOURCE:

Thiokol/Ventron Div., Danvers, MA, 01923, USA Precious Met. [Proc. Int. Precious Met. Inst. Conf.], SOURCE: 5th (1982), Meeting Date 1981, 3-10. Editor(s): Zysk,

Edward D. Pergamon: Willowdale, Ont.

CODEN: 48RYAT

DOCUMENT TYPE: Conference LANGUAGE: English

NaBH4 recovery operations are used in precious metal recovery applications. The most established precious metal recovery use is Ag removal from spent photog. fixing solns. Ag recovery of >99% at >95% purity can be achieved with this process. Applications have been developed for reduction of AgCl via a process which is capable of achieving extremely pure Ag with >99% recovery of Ag from scrap. Applications are being developed in the areas of Pt, Pd, and Rh

recovery from spent catalyst materials. NaBH4 is being used com. for

recovery of most of the precious metals.

IT 16940-66-2

RL: PROC (Process)

(in precious metal recovery)

RN

16940-66-2 HCAPLUS
Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME) CN

Na +

**7440-06-4P**, preparation **7440-22-4P**, preparation TΤ

RL: PUR (Purification or recovery); PREP (Preparation)

(recovery of, sodium borohydride in)

7440-06-4 HCAPLUS RN

Platinum (8CI, 9CI) (CA INDEX NAME) CN

Pt

7440-22-4 HCAPLUS RN

Silver (8CI, 9CI) (CA INDEX NAME) CN

Ag

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L33 ANSWER 35 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1982:168042 HCAPLUS
DOCUMENT NUMBER:
                        96:168042
                        Sodium borohydride controls heavy metal
TITLE:
                         discharge
AUTHOR (S):
                         Cook, Michael M.; Lander, Joseph A.
                         Ventron Div., Thiokol Corp., Danvers, MA, USA
CORPORATE SOURCE:
                         Pollution Engineering (1981), 13(12), 36-8
SOURCE:
                         CODEN: PLENBW; ISSN: 0032-3640
DOCUMENT TYPE:
                         Journal
                         English
LANGUAGE:
     Discharge of heavy metals into sewer systems and open waters is
     a major concern to both publicly owned treatment works and industries.
     Using NaBH4 as an alternative chemical treatment has enabled a number of plants
     to meet wastewater discharge limits in a cost-effective manner. It
     provides an effective and economical method for the removal of pollutants
     or for the recovery of precious metals, including Co, Cu, Au, Ir, Pb, Hg, Ni, Pd, Pt, Rh, and Hg. The metal
     is chemical reduced to the elemental state, usually as a compact precipitate
The
     NaBH4 process is irreversible, extremely rapid, and results in nearly
     total removal of the dissolved metals even at low initial concentration
     7439-92-1P, preparation 7440-02-0P, preparation
     7440-05-3P, preparation 7440-06-4P, preparation
     7440-22-4P, preparation 7440-48-4P, preparation
     7440-50-8P, preparation 7440-57-5P, preparation
     RL: PREP (Preparation)
        (recovery of, from wastewater, precipitation by sodium
        borohydride in)
     7439-92-1 HCAPLUS
RN
     Lead (8CI, 9CI) (CA INDEX NAME)
CN
Pb
     7440-02-0 HCAPLUS
RN
     Nickel (8CI, 9CI) (CA INDEX NAME)
CN
Νi
     7440-05-3 HCAPLUS
ВN
     Palladium (8CI, 9CI) (CA INDEX NAME)
CN
Pd
     7440-06-4 HCAPLUS
RN
     Platinum (8CI, 9CI) (CA INDEX NAME)
CN
Pt
     7440-22-4 HCAPLUS
RN
     Silver (8CI, 9CI) (CA INDEX NAME)
CN
```

Ag

RN 7440-48-4 HCAPLUS CN Cobalt (8CI, 9CI) (CA INDEX NAME)

Co

RN 7440-50-8 HCAPLUS CN Copper (7CI, 8CI, 9CI) (CA INDEX NAME)

Cu

RN 7440-57-5 HCAPLUS CN Gold (8CI, 9CI) (CA INDEX NAME)

Au

IT 16940-66-2

RL: PROC (Process)

(wastewater treatment by, metal precipitation in)

RN 16940-66-2 HCAPLUS

CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

● Na+

L33 ANSWER 36 OF 36 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1975:485606 HCAPLUS DOCUMENT NUMBER: 83:85606 Aromatic hydrogenation using sodium borohydride TITLE: reduced transition metal supported catalysts INVENTOR(S): Benson, Robert F. PATENT ASSIGNEE(S): Texaco Inc., USA U.S., 5 pp. SOURCE: CODEN: USXXAM DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION: KIND DATE APPLICATION NO. DATE PATENT NO. ---------<del>-----</del> -----US 1972-319574 19721229 US 1971-158581 A1 19710630 US 3869521 A 19750304 PRIORITY APPLN. INFO.: Supported transition metal catalysts are provided by a method comprising impregnating a catalyst support with a solution comprising a catalyst metal ion; at least partially drying the saturated catalyst support; admixing the dried catalyst support with sodium borohydride aqueous solution until the evolution of gas ceases, thereby obtaining an active supported metal catalyst; and thereafter, recovering the active catalyst. IT 7440-02-0, uses and miscellaneous RL: CAT (Catalyst use); USES (Uses) (catalysts, for demethylation of methylnaphthalene, manufacture of) RN7440-02-0 HCAPLUS Nickel (8CI, 9CI) (CA INDEX NAME) CN Ni 7440-06-4, uses and miscellaneous IT RL: CAT (Catalyst use); USES (Uses) (catalysts, for hydrogenation of benzene, manufacture of) RN7440-06-4 HCAPLUS CN Platinum (8CI, 9CI) (CA INDEX NAME) Pt TТ 7440-05-3, uses and miscellaneous RL: CAT (Catalyst use); USES (Uses) (catalysts, for hydrogenation of toluene, manufacture of) RN 7440-05-3 HCAPLUS CN Palladium (8CI, 9CI) (CA INDEX NAME) Pd IT 16940-66-2 RL: RCT (Reactant); RACT (Reactant or reagent) (reduction by, in preparation of hydrogenation catalysts) 16940-66-2 HCAPLUS RN CN Borate(1-), tetrahydro-, sodium (8CI, 9CI) (CA INDEX NAME)

● Na+

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=> => d stat que 153
            795 SEA FILE=REGISTRY ABB=ON PLU=ON TETRAHYDROBORATE OR HYDROBORA
L1
            11 SEA FILE=REGISTRY ABB=ON PLU=ON SODIUM BOROHYDRATE?/CN OR
L2
               LITHIUM BOROHYDR?/CN OR POTASSIUM BOROHYDR?/CN
L3
         104214 SEA FILE=REGISTRY ABB=ON PLU=ON METAL OR METALS OR MAGNESIUM
        128483 SEA FILE=REGISTRY ABB=ON PLU=ON BORATE
L4
         15523 SEA FILE=HCAPLUS ABB=ON PLU=ON L1 OR TETRAHYDROBORATE OR
L5
               HYDROBORATE
                                              52 TERMS
L6
               SEL PLU=ON L2 1- CHEM :
L7
         15422 SEA FILE=HCAPLUS ABB=ON PLU=ON L6
         15583 SEA FILE=HCAPLUS ABB=ON PLU=ON L7 OR (SODIUM OR LITHIUM OR
L8
               POTASSIUM) (2A) BOROHYDR?
       4137887 SEA FILE=HCAPLUS ABB=ON PLU=ON L3 OR METAL OR MAGNESIUM
L9
        166954 SEA FILE=HCAPLUS ABB=ON PLU=ON L4 OR BORATE
L10
         15254 SEA FILE=HCAPLUS ABB=ON PLU=ON L10(L)L9
L12
         38473 SEA FILE=HCAPLUS ABB=ON PLU=ON L10(L)(RCT/RL OR RACT/RL)
L14
         250013 SEA FILE=HCAPLUS ABB=ON PLU=ON L9(L) (RCT/RL OR RACT/RL)
L15
          2579 SEA FILE=HCAPLUS ABB=ON PLU=ON L12 AND L14
L16
          1952 SEA FILE=HCAPLUS ABB=ON PLU=ON L16 AND L15
L17
         104576 SEA FILE=REGISTRY ABB=ON PLU=ON PALLADIUM/BI
L18
           208 SEA FILE=REGISTRY ABB=ON PLU=ON PT/MF
L19
        513099 SEA FILE=HCAPLUS ABB=ON PLU=ON L18 OR L19 OR PALLADIUM OR
L21
               PLATINUM OR PD OR PT
          3041 SEA FILE=HCAPLUS ABB=ON PLU=ON (L5 OR L8)(L)(PREP?/RL OR
L25
               PREP OR PREPARTION)
L26
            78 SEA FILE=HCAPLUS ABB=ON PLU=ON L17 AND L25
            11 SEA FILE=HCAPLUS ABB=ON PLU=ON L26 AND L21
L27
             9 SEA FILE=HCAPLUS ABB=ON PLU=ON L27 AND PD=<NOVEMBER 25, 2003
L28
L30
          524 SEA FILE=HCAPLUS ABB=ON PLU=ON (L5 OR L8)(L)(PURIFI?/RL OR
               PURIFI? OR RECOVER?)
L31
           152 SEA FILE=HCAPLUS ABB=ON PLU=ON L30 AND L10 AND L9
            38 SEA FILE=HCAPLUS ABB=ON PLU=ON
L32
                                               L31 AND L9 AND L21
            36 SEA FILE=HCAPLUS ABB=ON PLU=ON
L33
                                               L32 NOT (L27 OR L28)
           294 SEA FILE=HCAPLUS ABB=ON PLU=ON
L44
                                               "SUDA S"/AU OR "SUDA SEIJIRAU"
               /AU
          1522 SEA FILE=HCAPLUS ABB=ON PLU=ON LI Z ?/AU OR "LI ZHOUPENG"/AU
L45
            52 SEA FILE=HCAPLUS ABB=ON PLU=ON "IWASE Y"/AU OR ("IWASE
L46
               YASUTOSHI"/AU OR "IWASE YASUYOSHI"/AU)
L47
             8 SEA FILE=HCAPLUS ABB=ON PLU=ON ("MORIGASAKI N"/AU OR
               "MORIGASAKI NOBUTO"/AU)
             7 SEA FILE=HCAPLUS ABB=ON PLU=ON L47 NOT (L33 OR L27 OR L28)
L48
             5 SEA FILE=HCAPLUS ABB=ON PLU=ON L44 AND L45 AND L46
L49
            44 SEA FILE=HCAPLUS ABB=ON PLU=ON L44 AND (L45 OR L46)
L50
            5 SEA FILE=HCAPLUS ABB=ON PLU=ON L45 AND L46
L51
            46 SEA FILE=HCAPLUS ABB=ON PLU=ON L48 OR L49 OR L50 OR L51
L52
            46 SEA FILE=HCAPLUS ABB=ON PLU=ON L52 NOT (L33 OR L27 OR L28)
L53
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<sup>=&</sup>gt; d ibib abs 153 1-46

L53 ANSWER 1 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:1249878 HCAPLUS

DOCUMENT NUMBER: 144:276900

TITLE: Development of the direct borohydride fuel cell

AUTHOR(S): Li, Z. P.; Liu, B. H.; Arai, K.; Suda,

S.

CORPORATE SOURCE: Materials & Energy Research Institute Tokyo, Ltd.,

Chino-shi, Nagano, 391-0301, Japan

SOURCE: Journal of Alloys and Compounds (2005), 404-406,

648-652

CODEN: JALCEU; ISSN: 0925-8388

PUBLISHER: Elsevier B.V.

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

AB A review on development of the direct borohydride fuel cells. The theor. energy conversion efficiency of the direct borohydride fuel cell (0.91) is larger than that of the polymer electrolyte membrane fuel cell (0.83). Recently, the cell power d. 290 mW/cm2 has been achieved. A five-cell stack with an effective area of 67 cm2 has demonstrated power of 110 W when the operation temperature is 60°, even if the stack operation

started at room temperature without humidification.

REFERENCE COUNT: 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 2 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:1249877 HCAPLUS

TITLE: Production of sodium borohydride by using dynamic

behaviors of protide at the extreme surface of

magnesium particles

AUTHOR(S): Suda, S.; Morigasaki, N.;

Iwase, Y.; Li, Z. P.

CORPORATE SOURCE: Department of Environmental and Chemical Engineering,

Chemical Energy Laboratory, Kogakuin University, 2665-1, Nakano-machi, Hachioji-shi, Tokyo, 192-0015,

Japan

SOURCE: Journal of Alloys and Compounds (2005), 404-406,

643-647

CODEN: JALCEU; ISSN: 0925-8388

PUBLISHER: Elsevier B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

AB An advanced process for the production of sodium borohydride (NaBH4) as a hydrogen storage material was developed, which applied the dynamic hydriding and dehydriding behaviors of protide (H-) in Mg-H system under transitional temperature conditions. An abundant natural resource named borax (Na2B4O7·10H2O) and the anhydrous sodium metaborate (NaBO2) recovered from the "spent fuel" as NaBO2.4H2O were used as the starting material in the present process. Powder-state Mg played an important role in the transitional hydriding and dehydriding process where the gaseous hydrogen was converted to protide at the extreme surface of Mg to form NaBH4 in exchange with the simultaneous transition of oxygen in NaBO2 to form MgO. In the present process, the protide as the most reactive state among the four states of hydrogen is applied for the synthesis of NaBH4, which can exist in metal-hydrogen complexes, such as NaAH4 and NaBH4. The NaBH4 yield was reached higher than 90% by a single batch process but was found to be largely dependent on the rate of temperature change and the particle

i.e., the sp. surface area of Mg particles.

REFERENCE COUNT: 31 THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Page 99

L53 ANSWER 3 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:491812 HCAPLUS

DOCUMENT NUMBER: 143:176139

AUTHOR (S):

TITLE: Performance improvement of a micro borohydride fuel

cell operating at ambient conditions Liu, B. H.; Li, Z. P.; Arai, K.; Suda,

s.

CORPORATE SOURCE: Department of Environmental and Chemical Engineering,

Kogakuin University, Hachioji, Tokyo, 192-0015, Japan

SOURCE: Electrochimica Acta (2005), 50(18), 3719-3725

CODEN: ELCAAV; ISSN: 0013-4686

PUBLISHER: Elsevier B.V. DOCUMENT TYPE: Journal

LANGUAGE: Journal English

AB In this study, aqueous borohydride solns. were employed to fuel a micro cell. Electrochem. performance of the micro borohydride fuel cell was tested at ambient conditions without any auxiliary facilities. Electrochem. impedance spectroscopy analyses were performed to characterize the cell performance. Both anion and cation exchange membranes were tried to sep. the fuel from the cathode. Membrane properties were found to be a decisive factor for cell performance. A maximum power d. of 40 mW/cm2 at room temperature was achieved when the Nafion NRE211 membrane was used. Hydrogen evolution at the anode side resulting from the competitive hydrolysis reaction influenced cell performance by preventing transfer of the electrolyte. The cell also demonstrated promising performance even when a silver cathode was used.

REFERENCE COUNT: 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 4 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

2004:1060839 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

142:25940

TITLE:

Production of alkali metal tetrahydroborates by

hydrogenation of alkali metal metaborates via alkaline

earth metal hydride intermediates

INVENTOR (S): Suda, Seijirau; Li, Zhoupeng;

Iwase, Yasuyoshi; Morigasaki, Nobuto

PATENT ASSIGNEE(S):

Japan

SOURCE:

U.S. Pat. Appl. Publ., 9 pp., Cont.-in-part of U.S.

Ser. No. 558,828.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2004249215	A1	20041209	US 2003-721479	20031125
JP 2004224684	A2	20040812	JP 2003-72466	20030317
PRIORITY APPLN. INFO.:			. US 2000-558828	A2 20000426
			JP 2002-344426	A 20021127
			JP 2003-72466	A 20030317

Alkali metal tetrahydroborates are produced in high yield by reaction of a AB suitable alkali metal borate (preferably an alkali metal metaborate) with a finely ground alkaline earth metal (preferably magnesium, ground to average particle diameter ≤100 µm) in the presence of H2 at below the equilibrium pressure for stable alkaline earth metal hydride formation. solid

reactants are mixed with hydrogen at ≤450°, and heated to 500-650°. Coke-oven gas is a suitable hydrogen source. Alkali metal tetrahydroborates have been suggested as a convenient source of fuel-cell-grade hydrogen.

L53 ANSWER 5 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:899747 HCAPLUS

DOCUMENT NUMBER: 143:175994

Sodium hydroborate as a hydrogen storing material TITLE:

Suda, Seijirou; Morigasaki, Nobuto; Iwase, AUTHOR (S):

Yasuyoshi; Li. Zhou Peng

Dep. of Engineering, Kogakuin University, Japan CORPORATE SOURCE:

Kagaku Sochi (2004), 46(10), 64-71 CODEN: KASOB7; ISSN: 0368-4849 SOURCE:

Kogyo Chosakai PUBLISHER:

DOCUMENT TYPE: Journal; General Review

LANGUAGE: Japanese

AB A review with 0 refs.

L53 ANSWER 6 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:446908 HCAPLUS

DOCUMENT NUMBER: 140:409212

TITLE: Method for producing tetrahydroborates

INVENTOR(S): Suda, Seijirau; Li, Zhoupeng;

Iwase, Yasuyoshi; Morigasaki, Nobuto

PATENT ASSIGNEE(S): Materials & Energy Research Institute Tokyo, Ltd.,

Japan

SOURCE: Eur. Pat. Appl., 11 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.		KIND	DATE	APPLICATION NO.	DATE
EP 1424310		A2	20040602	EP 2003-27149	20031126
R: AT,	BE, CH,	DE, DK	, ES, FR,	GB, GR, IT, LI, LU,	NL, SE, MC, PT,
IE,	SI, LT,	LV, FI	, RO, MK,	CY, AL, TR, BG, CZ,	EE, HU, SK
JP 20042246	84	A2	20040812	JP 2003-72466	20030317
PRIORITY APPLN.	INFO.:			JP 2002-344426	A 20021127
				JP 2003-72466	A 20030317

JP 2003-72466 A 20030317

AB Tetrahydroborates, such as NaBH4, LiBH4, and KBH4, are prepared by reacting a powdery mixture containing a borate and magnesium in a hydrogen atmospheric under

pressure at 500-650°. The hydrogen source can be coke oven gas. The reaction can be carried out in the presence of a hydrogenation catalyst. The starting materials have an average particle diameter of < 100  $\mu m_{\odot}$ 

L53 ANSWER 7 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:306656 HCAPLUS

DOCUMENT NUMBER: 141:210022

TITLE: Metal-hydrogen complex compounds as hydrogen storage

materials

AUTHOR(S): Suda, S.; Iwase, Y.;

Morigasaki, N.; Li, Z.-P.

CORPORATE SOURCE: Department of Environmental & Chemical Engineering,

Kogakuin University, Hachioji, 192-0015, Japan

SOURCE: Advanced Materials for Energy Conversion II,

Proceedings of a Symposium held during the TMS Annual Meeting, 2nd, Charlotte, NC, United States, Mar. 14-18, 2004 (2004), Meeting Date 2004, 123-133. Editor(s): Chandra, Dhanesh; Bautista, Renato G.; Schlapbach, Lous. Minerals, Metals & Materials

Society: Warrendale, Pa.

CODEN: 69FHAF; ISBN: 0-87339-574-3

DOCUMENT TYPE: Conference LANGUAGE: English

AB A new production process for NaBH4 as H storage material was developed by

using NaBO2, according to NaBO2 + 2Mg + 2H2 → NaBH4 + 2MgO at a

fast reaction rate by changing the temperature of the system dynamically. T process uses the transitional states of H at the surface of Mg when it is hydrogenated and dehydrogenated at a constant rate of temperature change,

dT/dt.

The rate of conversion is dependent on dT/dt and is regulated by the extent of oxidation of the Mg particles. The role of H- in Mg·(2H-) under transient conditions, Mg + H2  $\rightarrow$  MgH2  $\rightarrow$  Mg

 $\cdot\,(2H\text{-})\,,$  is discussed with regard to the exchange reaction between NaBO2 and NaBH4.

REFERENCE COUNT:

30 THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 8 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2004:83205 HCAPLUS

DOCUMENT NUMBER:

140:256177

TITLE:

Evaluation of alkaline borohydride solutions as the

fuel for fuel cell

AUTHOR (S):

Li, Z. P.; Liu, B. H.; Arai, K.; Asaba, K.;

Suda, S.

CORPORATE SOURCE:

Department of Environmental & Chemical Engineering, Kogakuin University, Hachioji-shi, Tokyo, 192-0015,

Japan

SOURCE:

Journal of Power Sources (2004), 126(1-2), 28-33

CODEN: JPSODZ; ISSN: 0378-7753

Elsevier Science B.V.

DOCUMENT TYPE:

Journal

LANGUAGE:

PUBLISHER:

English

A fuel cell was assembled using alkaline borohydride solns. as the fuel. The sp. gr., sp. viscosity, and m.ps. of the borohydride solns. were evaluated. With increasing the NaBH4 concentration, the sp. gr. was decreased but the viscosity was increased consistently. The effects of NaBH4 and NaOH concns. on the cell and electrode polarizations were investigated. The increase of NaBH4 concentration improved the anode performance only a little

but significantly increased the cathode polarization due to the BH4crossover. As a result, the cell polarization increased when using highly concentrated borohydride solns. Based on the Nernst's equation, the increase

of

NaOH concentration is favorable for the anode, but detrimental to the cathode polarization. In principle, the cell polarization has nothing to do with the NaOH concentration according to the cell reaction NaBH4 + O2 = NaBO2 + 2H2O.

However, the exptl. results showed that the high concentration of NaOH affected the cell polarization at higher current densities.

REFERENCE COUNT:

THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS 12 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 9 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:601030 HCAPLUS

DOCUMENT NUMBER: 139:309993

TITLE: Protide compounds in hydrogen storage systems AUTHOR(S): Li, Z. P.; Liu, B. H.; Arai, K.; Morigazaki,

N.; Suda, S.

CORPORATE SOURCE: Department of Environmental and Chemical Engineering,

Kogakuin University, Hachioji-shi, Tokyo, 192-0015,

Japan

SOURCE: Journal of Alloys and Compounds (2003), 356-357,

469-474

CODEN: JALCEU; ISSN: 0925-8388

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

AB Based on 3 chemical states of hydrogen, H-, HO and H+, a triangular H energy

system is proposed. The energy transfer between the 3 states is

discussed. Expts. prove that NaBH4 as a H- carrier can release its energy

directly through a borohydride fuel cell or generate H2 for polymer electrolyte membrane fuel cells. The used fuel, Na meta-borate, can be

converted back to NaBH4 through a reaction with MgH2.

REFERENCE COUNT: 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 10 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:458475 HCAPLUS

DOCUMENT NUMBER: 139:216877

TITLE: A Fuel Cell Development for Using Borohydrides as the

Fuel

AUTHOR(S): Li, Z. P.; Liu, B. H.; Arai, K.; Suda,

S.

CORPORATE SOURCE: Department of Environmental and Chemical Engineering,

Kogakuin University, Nakano-machi 2665-1,

Hachioji-shi, Tokyo, 192-0015, Japan

SOURCE: Journal of the Electrochemical Society (2003), 150(7),

A868-A872

CODEN: JESOAN; ISSN: 0013-4651

PUBLISHER: Electrochemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

AB A fuel cell was developed using borohydride solns. as the fuel. The cell consisted of an anode made of a Zr-Ni alloy, a cathode made of Pt/C, and a Na+ form Nafion membrane as the electrolyte. The borohydride-fueled cell showed an open-circuit voltage of 1.3 V, compared with 1.0 V for a hydrogen gas-fueled one. The anode exhibited a small polarization property compared with the cathode. The cathode polarization was the main reason for the cell voltage drop with increasing currents. When a Nafion membrane was used as the electrolyte, cations (Na+) were the charge carrier in it. Compared with Nafion 112 membrane, Nafion 117 membrane demonstrated a considerable resistance to borohydride crossover and resulted in acceptable cell performance. However, there are several problems such as H2 evolution during operation, BH-4 crossover, NaOH accumulation at the cathode, and NaBO2 accumulation at the anode in recent systems. Further effort is needed to develop the fuel cell using borohydrides as the fuel.

REFERENCE COUNT: 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 11 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

2003:324965 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 139.103130

Preparation of potassium borohydride by a TITLE:

mechano-chemical reaction of saline hydrides with

dehydrated borate through ball milling

AUTHOR (S): Li, Zhou Peng; Liu, Bin Hong; Morigasaki,

Nobuto; Suda, Sejirau

Department of Environmental and Chemical Engineering, CORPORATE SOURCE:

Kogakuin University, Hachioji-shi, Tokyo, 192-0015,

Japan

SOURCE: Journal of Alloys and Compounds (2003), 354(1-2),

243-247

CODEN: JALCEU; ISSN: 0925-8388

Elsevier Science B.V. PUBLISHER:

DOCUMENT TYPE: Journal English LANGUAGE:

A convenient method was developed to synthesize potassium borohydride by a mechanochem. reaction of saline hydrides with dehydrated borates using a planetary ball mill. Among the tested saline hydrides, MgH2 was the most effective reactant for borohydride formation. In order to improve the borohydride yield, it was necessary to add an excess amount of MgH2. When a 35% excess of MgH2 was added, potassium borohydride yield reached 100%. The borohydride formation was strongly dependent on the water content in reactants. When the water content in KBO2 samples was >24.8 weight%

(equivalent

to KBO2·1.5H2O), no borate was converted into borohydride by the mechanochem. reaction.

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 8

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 12 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2003:99330 HCAPLUS

DOCUMENT NUMBER:

138:371547

TITLE:

A study on recycle of borohydride liquid fuel

AUTHOR (S):

Morigasaki, Nobuto; Tanisawa, Kazuhiro;

CORPORATE SOURCE:

Li, Zhoupeng; Suda, Seijirau

Graduate School of Applied Chemistry and Chemical Engineering, Kogakuin University, Tokyo, Japan

SOURCE:

Kogakuin Daigaku Kenkyu Hokoku (2002), 93, 55-59 CODEN: KDKHAY; ISSN: 0368-5098

Kogakuin Daigaku

PUBLISHER: DOCUMENT TYPE:

Journal

LANGUAGE:

Japanese

Borohydrides can be formed by the reaction of anhydrous metaborate with Mq hydride. However, the borohydride conversion rate was influenced by the metaborate source. Anhydrous sodium metaborate can be produced by dehydrating the NaBO2·4H2O crystals from used fuel or the commercialized NaBO2·4H2O. The NaBH4 conversion rate based on NaBO2 from used fuel was lower than that based on the anhydrous borate form the commercialized NaBO2·4H2O. It was because the sample obtained from used fuel was not completely dehydrated.

L53 ANSWER 13 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:99329 HCAPLUS

DOCUMENT NUMBER: 138:371546

TITLE: The electric discharge mechanism of the liquid type

fuel cell using borohydride complex ion

AUTHOR(S): Arai, Kunihito; Chiku, Satoshi; Li, Zhoupeng

; Suda, Seijirau

CORPORATE SOURCE: Materials & Energy Research Institute Tokyo, Ltd.,

Japan

SOURCE: Kogakuin Daigaku Kenkyu Hokoku (2002), 93, 49-53

CODEN: KDKHAY; ISSN: 0368-5098

PUBLISHER: Kogakuin Daigaku

DOCUMENT TYPE: Journal LANGUAGE: Japanese

In this liquid type fuel cell using borohydride complex ion (called "BFC"), it was found that alkaline type reactions had occurred at each electrode, though this fuel cell used the pos. ion exchange membrane (i.e. an acid membrane) as the electrolyte. These reactions are involved in the fact that K+ ion which exists so much at an anode side moved in the electrolyte as an elec. charge carrier. Moreover, when we used the fluorinated metal hydride as an anode catalyst, we obtained potential as much as Pd.

L53 ANSWER 14 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:99328 HCAPLUS

DOCUMENT NUMBER: 138:355708

TITLE: A review on the methods of synthesis of alkali metal

borohydrides

AUTHOR(S): Iwase, Yasuyoshi; Li, Zhoupeng;

Suda, Seijirau

CORPORATE SOURCE: Materials & Energy Research Institute Tokyo, Ltd.,

Japan

SOURCE: Kogakuin Daiqaku Kenkyu Hokoku (2002), 93, 43-48

CODEN: KDKHAY; ISSN: 0368-5098

PUBLISHER: Kogakuin Daigaku

DOCUMENT TYPE: Journal; General Review

LANGUAGE: Japanese

AB A review of articles and patents on the synthesis methods of alkali metal

borohydrides to explore the required conditions for new and more

economical methods of producing the alkali metal borohydrides. Based on this overview and our experiment results, we determined the conditions to

synthesize

borohydrides (BH4-) from borate salts: (a) reaction proceeds under anhydrous condition or in aprotic solvent and (b) borate salts react with other protide compds. (H-) to form BH4-, but never form BH4- with proton.

Page 111

L53 ANSWER 15 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2002:972505 HCAPLUS

DOCUMENT NUMBER:

138:347880

TITLE:

Preparation of sodium borohydride by the reaction of MgH2 with dehydrated borax through ball milling at

room temperature

AUTHOR (S):

Li, Z. P.; Morigazaki, N.; Liu, B. H.;

Suda, S.

CORPORATE SOURCE:

Department of Environmental and Chemical Engineering, Kogakuin University, Hachioji-shi, Tokyo, 192-0015,

Japan

SOURCE:

Journal of Alloys and Compounds (2003), 349(1-2),

232-236

CODEN: JALCEU; ISSN: 0925-8388

PUBLISHER:

Elsevier Science B.V.

DOCUMENT TYPE:

Journal

English

LANGUAGE:

A convenient method was developed to synthesize NaBH4 by the reaction of MqH2 with Na2B4O7 through ball milling at room temperature To improve the Na

borohydride yield, Na compds. were added to compensate the Na

insufficiency in reactants when MgH2 instead of NaH was used as the

reducing agent. Na2CO3 addition was better than NaOH or Na2O2 addition in increasing the borohydride yield.

REFERENCE COUNT:

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 16 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2002:95069 HCAPLUS

DOCUMENT NUMBER: 136:312397

TITLE: Effects of surface structure of fluorinated AB2 alloys

on their electrodes and battery performances

AUTHOR(S): Li, Z. P.; Liu, B. H.; Hitaka, K.;

Suda, S.

CORPORATE SOURCE: Department of Environmental and Chemical Engineering,

Kogakuin University, Hachioji, 192-0015, Japan

SOURCE: Journal of Alloys and Compounds (2002), 330-332,

776-781

CODEN: JALCEU; ISSN: 0925-8388

PUBLISHER: Elsevier Science S.A.

DOCUMENT TYPE: Journal LANGUAGE: English

AB Fluorination technique effectively improve the activation property and rate discharge capabilities of AB2 alloys. In this paper, the performance of sealed batteries using fluorinated AB2 alloys were investigated. Compared with the sealed battery using untreated AB2 alloys, those using fluorinated AB2 alloys showed better activation property, longer cycle life and higher rate discharge capacity. The rate discharge capacity of the sub-C sealed battery using duplicated fluorination treated AB2 alloys reached up to 2.3 AH at 4C (10.4A). The batteries using fluorinated AB2 alloys sustained over 300 cycles at 0.2C rate, but its discharge capacity

degraded quickly when being cycled at 1C rate.

REFERENCE COUNT: 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Page 113

L53 ANSWER 17 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:300367 HCAPLUS

DOCUMENT NUMBER: 135:95104

TITLE: Fluorinated metal hydrides for the catalytic

hydrolysis of metal-hydrogen complexes

AUTHOR(S): Suda, S.; Sun, Y.-M.; Uchida, M.; Liu,

B.-H.; Mormitu, S.; Arai, K.; Zhou, Y.; Tsukamoto, N.;

Candra, Y.; Li, Z.-P.

CORPORATE SOURCE: Chemical Energy Laboratory, Kogakuin University,

Tokyo, 192-0015, Japan

SOURCE: Metals and Materials International (2001), 7(1), 73-75

CODEN: MMIECY

PUBLISHER: Korean Institute of Metals and Materials

DOCUMENT TYPE: Journal LANGUAGE: English

This paper is related to the hydrogen storage (H-storage) system composed of an alkaline solution that contains metal-hydrogen complex ions. The BH4--complex ions stabilized when dissolved in aqueous KOH or NaOH to form homogeneous solns. However, it is necessary to apply an appropriate catalyst for a high rate generation of hydrogen under atmospheric pressure and temperature conditions. The authors intended to develop catalysts suitable for the hydrolysis of BH4 -complex ion solns. and found that Mg2Ni exhibits excellent catalytic functions in the hydrolysis, in particular, after the fluorination treatment (F-treatment). In this work, the effects of F-treatment on the rates of hydrolysis were evaluated for the hydrided and unhydrided Mg2Ni. No hydrogenation in addition to F-treatment is required in view of hydrolysis kinetics and cost-effectiveness.

REFERENCE COUNT: 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 18 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:206623 HCAPLUS

DOCUMENT NUMBER: 134:313607

TITLE: Catalytic generation of hydrogen by applying

fluorinated-metal hydrides as catalysts

Suda, S.; Sun, Y.-M.; Liu, B.-H.; Zhou, Y.; AUTHOR (S):

Morimitsu, S.; Arai, K.; Tsukamoto, N.; Uchida, M.;

Candra, Y.; Li, Z.-P.

CORPORATE SOURCE: Chemical Energy Laboratory, Department of

Environmental and Chemical Engineering, Kogakuin

University, Hachioji, 192-0015, Japan

SOURCE: Applied Physics A: Materials Science & Processing

(2001), 72(2), 209-212 CODEN: APAMFC; ISSN: 0947-8396

PUBLISHER: Springer-Verlag

39

DOCUMENT TYPE: Journal LANGUAGE: English

Metal-hydrogen complexes such as NaAlH4, KBH4, and NaBH4 are known as high H-content materials. The highly reactive natures of these materials against moist air and water can be easily stabilized in aqueous KOH and NaOH solns. Accordingly, it is required to develop catalysts suitable for generating hydrogen from the stabilized metal-hydrogen complexes in alkaline solns. This work is aimed at developing catalysts that can generate hydrogen from such solns. with considerably high kinetics under moderate temperature and pressure conditions. We have found that Mg2Ni, a typical high-temperature hydriding alloy, exhibits excellent functions as a catalyst

for

the hydrolysis of BH4--ion-containing solns. The fluorination-treatment (F-treatment) effects on granular particles of Mg2Ni and Mg2NiH4 are reported in this paper.

REFERENCE COUNT:

THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 19 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:88882 HCAPLUS

DOCUMENT NUMBER:

134:298330

TITLE:

Effects of fluorination of AB2-type alloys and of

mixing with AB5-type alloys on the charge-discharge

characteristics

AUTHOR (S):

Higuchi, E.; Toyoda, E.; Li, Z. P.;

CORPORATE SOURCE:

Suda, S.; Inoue, H.; Nohara, S.; Iwakura, C. Faculty of Engineering, Department of Environmental

and Chemical Engineering, Chemical Energy Lab, Koqakuin University, Hachioji-shi, Tokyo, 192-0015,

Japan

SOURCE:

Electrochimica Acta (2001), 46(8), 1191-1194

CODEN: ELCAAV; ISSN: 0013-4686

PUBLISHER:

Elsevier Science Ltd.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

The effects of fluorination (F-treatment) of AB2-type Laves-phase Zr0.9Ti0.1V0.2Mn0.6Co0.1Ni1.1 (AB2) alloys and of mixing with untreated AB5-type MmNi3.68Co0.78Mn0.36Al0.28 (AB5) alloys on their charge-discharge characteristics were investigated. The F-treatment of the AB2 alloy greatly increased initial activation rate and discharge capacity and slightly improved high-rate dischargeability. The mixing of the F-treated AB2 alloy with untreated AB5 (<25  $\mu$ m) alloy decreased somewhat the maximum discharge capacity compared to the fluorinated AB2 alloy but it greatly improved the high-rate dischargeability. In particular, the high-rate dischargeability for the AB2(F)-AB5 (<25  $\mu$ m) mixture pos. deviated from the estimated values based on the maximum discharge capacity for both alloys

and

their content, suggesting that the AB5 alloy can work as a catalyst for dehydriding the AB2 alloy hydride besides as a hydrogen reservoir; a synergistic effect appeared by the mixing.

REFERENCE COUNT:

17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 20 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:147662 HCAPLUS

DOCUMENT NUMBER: 132:200265

TITLE: Effects of fluorination temperature on surface

structure and electrochemical properties of AB2

electrode alloys

AUTHOR(S): Li, Z. P.; Higuchi, E.; Liu, B. H.;

Suda, S.

CORPORATE SOURCE: Department of Environmental and Chemical Engineering,

Kogakuin University, Hachioji, 192-0015, Japan Electrochimica Acta (2000), 45(11), 1773-1779

SOURCE: Electrochimica Acta (2000), 45(11), 177
CODEN: ELCAAV; ISSN: 0013-4686

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

AB In order to accelerate the activation of AB2 electrode alloys, a new fluorination treatment has been developed which removes oxides from alloy surfaces and implants metallic Ni into the alloy surface layer. The behavior of Ni during fluorination treatment depends strongly on the treatment temperature. The constituent Ni of the alloy dissolves into the treatment solution at a low temperature, but Ni complex ion in treatment solution is

chemical reduced by constituent elements at a high temperature. The temperature is the

key factor of the fluorination treatment. The electrochem. properties were improved through increasing the treatment temperature due to the Ni implantation into surface layer.

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 21 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2000:6032 HCAPLUS

DOCUMENT NUMBER:

132:95674

TITLE:

Electrochemical properties and characteristics of a

fluorinated AB2-alloy

AUTHOR (S):

Li, Z. P.; Higuchi, E.; Liu, B. H.;

Suda, S.

CORPORATE SOURCE:

Department of Environmental & Chemical Engineering, Kogakuin University, Hachioji, 192-0015, Japan

SOURCE:

Journal of Alloys and Compounds (1999), 293-295,

593-600

CODEN: JALCEU; ISSN: 0925-8388

PUBLISHER:

Elsevier Science S.A.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

A series of fluorination techniques have been developed in this laboratory to improve the surface condition of AB2 alloy. The fluorination was found effective for removing oxide layer and implanting catalytic Ni to the alloy surfaces. The developed fluorination techniques improved considerably the initial activation characteristics, decreased the electrochem. reaction impedance and increased the rate capacity of AB2 electrodes.

REFERENCE COUNT:

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 22 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1999:482282 HCAPLUS

DOCUMENT NUMBER: 131:274162

TITLE: Fluorinated Laves-AB2 alloy as the negative electrode

in Ni-MH battery

AUTHOR(S): Higuchi, Eiji; Miyoshi, Hirokazu; Sugimoto, Hiroyuki;

Li, Zhoupeng; Suda, Seijirau

CORPORATE SOURCE: Grad. Sch. Ind. Chem., Kogakuin Univ., Tokyo,

163-8677, Japan

SOURCE: Kogakuin Daigaku Kenkyu Hokoku (1999), 86, 69-75

CODEN: KDKHAY; ISSN: 0368-5098

PUBLISHER: Kogakuin Daigaku

DOCUMENT TYPE: Journal LANGUAGE: Japanese

AB AB5 hydriding alloys as neg. electrode materials have been well developed for Ni-MH battery. Many scientists have paid attention to AB2 alloy due to the high-capacity of AB2 alloy compared with AB5 alloy. However, AB2 alloy usually showed poor activation and kinetic properties. It is possible to overcome these shortcomings by surface modification through fluorination. Aiming for practical use, we investigated the performances of practical electrode of fluorinated AB2 alloy to improve the rate of discharge capability and cycle life. It is found that the connection of lead line to electrode conductor of the electrode, and the pore size of the Ni foam exerted the big influence on electrode properties.

L53 ANSWER 23 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1999:482280 HCAPLUS

DOCUMENT NUMBER: 131:274160

TITLE: Effects of the fluorination treatment on the surface

structure and electrochemical properties of Zr-based

AB2-type hydriding alloys

AUTHOR(S): Okutsu, Akira; Nakamura, Kazuhiro; Li,

Zhoupeng; Suda, Seijirau

CORPORATE SOURCE: Grad. Sch. Ind. Chem., Kogakuin Univ., Tokyo,

163-8677, Japan

SOURCE: Koqakuin Daiqaku Kenkyu Hokoku (1999), 86, 57-62

CODEN: KDKHAY; ISSN: 0368-5098

PUBLISHER: Kogakuin Daigaku

DOCUMENT TYPE: Journal LANGUAGE: Japanese

AB AB5 alloys have been used in Ni-MH battery as neg. electrode materials. In this work, the activation and reactivity of Zr-based AB2 alloys whose capacities were larger than AB5 alloys, have been improved by fluorination. The effects of fluorination technique combining with plenary ball milling on electrochem. properties have been investigated to improve performances of AB2 alloys for developing the high power secondary battery of elec. vehicle. It was found that oxides can be removed from alloy surfaces while metallic Ni can be implanted to alloy surfaces so that the sp. surface area of the fluorinated AB2 alloy was increased. The electrochem. activation, high rate charge and discharge capability were improved through surface modification by fluorination.

L53 ANSWER 24 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1998:544811 HCAPLUS

DOCUMENT NUMBER:

129:163814

TITLE:

Surface passivation of metal hydrides for applications

AUTHOR(S): Suda, S.; Li, Z.-P.; Sun, Y.-M.;

Liu, B.-H.; Gao, X.-P.

CORPORATE SOURCE:

Department of Environmental & Chemical Engineering,

Kogakuin University, Tokyo, 192-0015, Japan

SOURCE:

Materials Research Society Symposium Proceedings (1998), 513 (Hydrogen in Semiconductors and Metals),

25-36

CODEN: MRSPDH; ISSN: 0272-9172

PUBLISHER:
DOCUMENT TYPE:

Journal; General Review

Materials Research Society

LANGUAGE:

English

34

AB A review with many refs. Properties and characteristics of hydriding alloys are strongly dependent on surface compns. and morphologies. For instance, oxides such as La2O3 on AB5 alloys and ZrO2 on AB2, AB, and body-centered-cubic (BCC) alloys act as the barriers for the conversion of mol. and ionic hydrogen to atomic hydrogen at the surface, thus reducing the kinetics in both the gas-solid and electrochem. reactions. Alloy surfaces chemical treated by an aqueous F-ion containing solution have been developed

to solve

such problems. F-treated surfaces exhibit significantly improved characteristics in regard to the hydrogen uptakes and the protection against impurities and electrolyte solution. In addition, highly conductive metallic Ni layers can be formed on the surface of the alloy particles by the fluorination. The authors report the properties and characteristics of fluorinated hydriding alloys, mainly of a typical AB2 Laves phase material which represents the difficult activation characteristics and poor long-term durability during electrochem. charge/discharge cycles.

REFERENCE COUNT:

THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 25 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1998:51782 HCAPLUS

DOCUMENT NUMBER: 128:117289

TITLE: The effects of fluorination solution composition on

the electrochemical properties of Zr0.9Ti0.1V0.2Mn0.6Co0.1Ni1.1 alloy

AUTHOR(S): Higuchi, Eiji; Sakasita, Masaya; Li, Zhoupeng

; Suda, Seijirau

CORPORATE SOURCE: Dep. Ind. Chem., Kogakuin Univ., Tokyo, 163-91, Japan

SOURCE: Kogakuin Daigaku Kenkyu Hokoku (1997), 83, 27-30

CODEN: KDKHAY; ISSN: 0368-5098

PUBLISHER: Kogakuin Daigaku

DOCUMENT TYPE: Journal LANGUAGE: Japanese

AB Effects of components of fluorination treatment solution on discharge properties of Zr-based AB2-type alloy Zr0.9Ti0.1V0.2Mn0.6Co0.1Ni1.1 were studied. It was found that Ni amount implanted on the hydriding alloy surface showed significant effects on the discharge capacity. The element dissoln. in the alloy during the treatment also showed similar effects.

L53 ANSWER 26 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1998:51781 HCAPLUS

DOCUMENT NUMBER:

128:117288

TITLE:

The effects of fluorination treatment on the electrochemical properties of Zr-based AB2-type

hydriding alloys

AUTHOR(S):

Okutsu, Akira; Sakasita, Masaya; Li, Zhoupeng

; Suda, Seijirau

CORPORATE SOURCE:

Dep. Ind. Chem., Kogakuin Univ., Tokyo, 163-91, Japan

SOURCE:

Kogakuin Daigaku Kenkyu Hokoku (1997), 83, 23-26

CODEN: KDKHAY; ISSN: 0368-5098

PUBLISHER:

Kogakuin Daigaku

DOCUMENT TYPE:

Journal

LANGUAGE:

Japanese

Fluorination treatment effects on discharge properties of Zr-based AB2-type hydriding alloys were studied. It was found that electrochem. activation properties could be improved and discharge capacity could be increased by implanting Ni on the surfaces of the hydriding alloys through the fluorination treatment.

L53 ANSWER 27 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

1997:456292 HCAPLUS ACCESSION NUMBER:

127:164304 DOCUMENT NUMBER:

Fluorination mechanism and its effects on the TITLE:

electrochemical properties of metal hydrides

Sakashita, M.; Li, Z. P.; Suda, S. AUTHOR (S):

CORPORATE SOURCE: Department of Chemical Engineering, Kogakuin

University, Nakano-machi 2665-1, Hachioji-shi, Tokyo,

192, Japan

Journal of Alloys and Compounds (1997), 253-254, SOURCE:

500-505

CODEN: JALCEU; ISSN: 0925-8388

PUBLISHER: Elsevier Journal DOCUMENT TYPE: English LANGUAGE:

Fluorination of hydriding alloys has been found effective for improving durability and initial discharge characteristics of hydride electrodes in Ni-MH rechargeable batteries. However, it has also been found that it decreases the discharge capacity because of the fluoride formed on the surface which decreases the elec. conductivity of the electrode. A more advanced

technique has been developed in this laboratory to implant metallic Ni in the fluoride layer to form a functionally graded surface layer. The proposed fluorination technique considerably improves the durability and initial activation characteristics of AB5-types of hydride, electrodes and also the initial activation characteristics of AB2 electrodes. The fluorination was found effective for removing the oxide layer which acts as an impedance to electrochem. hydrogen uptake.

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT:

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L53 ANSWER 28 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1996:430623 HCAPLUS

DOCUMENT NUMBER:

125:92484

TITLE:

An investigation of Ca-based hydride electrode

materials

AUTHOR (S):

Li, Z. P.; Matsuoka, T.; Suda, S.

CORPORATE SOURCE:

Dep. Chem. Eng., Kogakuin Univ., Tokyo, 192, Japan Vacuum (1996), 47(6-8, Proceedings of the 13th

SOURCE:

International Vacuum Congress and the 9th International Conference on Solid Surfaces, 1995),

893-897

CODEN: VACUAV; ISSN: 0042-207X

PUBLISHER:

Elsevier Journal

DOCUMENT TYPE: English LANGUAGE:

The effects of La-substitution for Ca in Ca1-xLaxNi5.7Al0.3 on metallog., structure and electrochem. properties have been investigated. It was found that Ca and Al segregated over the grain boundaries of the alloys. Based on La substituted CaNi5-type alloys, we studied the Zr and Ti addition effects as well. It was found that Zr-Ni phases or Ti-Ni phases deposited on the boundaries of AB5 phase improved the anti-corrosion property of the alloys. Annealing improves the electrochem. durability of the Cal-xLaxNi4.7Al0.3 alloys by reducing A2B7 phase and Ca segregation on the grin boundary.

L53 ANSWER 29 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:199099 HCAPLUS

DOCUMENT NUMBER: 124:265514

TITLE: A new family of hydride electrode materials based on

CaNi5-type alloys

AUTHOR(S): Matsuoka, Takashi; Li, Zhoupeng; Uchida,

Masaki; Suda, Seijirau

CORPORATE SOURCE: Kogakuin Univ., Tokyo, 163-91, Japan

SOURCE: Kogakuin Daigaku Kenkyu Hokoku (1995), 78, 85-7

CODEN: KDKHAY; ISSN: 0368-5098

PUBLISHER: Kogakuin Daigaku

DOCUMENT TYPE: Journal LANGUAGE: Japanese

AB Charge-discharge characteristic of Cal-xAxNi5-y-zMyNz (A = La, Ce, Mg, Sr; M = Sn, Si, Co; N = Al; 0≤x, y, z≤1) hydride electrode were examined The discharge capacity of the Ca-based alloys was 300-350 mAh/g at discharge current of 150 mA/g and was lower than that of CaNi5. In comparison to the M and/or N substitution, A substitution was effective on the improvement of cycling durability and kept the discharge capacity at a high level. The most durable electrodes were obtained with A = La and x = 0.6.

L53 ANSWER 30 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:106282 HCAPLUS

DOCUMENT NUMBER: 124:180993

TITLE: The Ni content and F-treatment effects on the

electrochemical cycling behavior of Zr0.5Ti0.5Nix

alloys

AUTHOR(S): Li, Zhoupeng; Takiguchi, Katsutoshi;

Suda, Seijirau

CORPORATE SOURCE: Kogakuin Univ., Tokyo, 163-91, Japan

SOURCE: Kogakuin Daigaku Kenkyu Hokoku (1995), 78, 81-4

CODEN: KDKHAY; ISSN: 0368-5098

PUBLISHER: Kogakuin Daigaku

DOCUMENT TYPE: Journal LANGUAGE: English

AB The alloys were prepared through arc melting of pure metals and pulverized to 26  $\mu m$  and the powders were treated with F ion-containing solution With increasing of Ni content in the alloy of x = 1.25-1.675, the discharge capacity increased and the electrochem. activation became easier, but with further addition of Ni discharge capacity decreased. The alloys of high content of Ni (x = 1.5-1.675) had higher discharge capacity but the alloys of lower capacity had better durability. F-treatment of alloys containing low Ni as x  $\leq$  1.25 decreased discharge capacity because of decrease of Ni content at the surface.

L53 ANSWER 31 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:106281 HCAPLUS

DOCUMENT NUMBER: 124:180992

TITLE: Effects of hydriding-dehydriding cycling on the P-C-T

and electrochemical properties of Cal-

χLaχNi4.7Al0.3 alloys

AUTHOR(S): Li, Zhoupeng; Suda, Seijirau

CORPORATE SOURCE: Kogakuin Univ., Tokyo, 163-91, Japan

SOURCE: Kogakuin Daigaku Kenkyu Hokoku (1995), 78, 77-80

CODEN: KDKHAY; ISSN: 0368-5098

PUBLISHER: Koqakuin Daigaku

DOCUMENT TYPE: Journal LANGUAGE: English

AB Electrodes were made through mixing the alloys of x = 0-0.6 with Ni and PTFE in weight ratio of 5:15:1 and through packing in Ni grid and pressing under 4 ton/cm2 to pellets of 13 mm dia. in 1.5 mm thickness. The cycling was done for composition of charge of 200 mA/g of 3 h and discharge of 150 mA/g in 6M KOH of 20° to 1.0 V vs. Ni/Ni hydroxide electrode. For XRD pellets of alloy/PTFE of 5/1 weight/weight were prepared After 40 cycles of hydriding/dehydriding at 40° no CaH2 signal was found. From P-C-T properties degradation of capacity was not observed after 50 cycles. La-substituted Ca-based alloys showed strong anti-disproportionation tendency to stabilize the lattice.

L53 ANSWER 32 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:71040 HCAPLUS

DOCUMENT NUMBER: 124:152479

TITLE: The annealing effect on the durabilities of La

substituted CaNi5-type alloys

AUTHOR(S): Li, Zhoupeng; Nagai, Susumu; Suda,

Seijirau

CORPORATE SOURCE: Kogakuin Univ., Tokyo, 163-91, Japan

SOURCE: Kogakuin Daigaku Kenkyu Hokoku (1995), 78, 71-5

CODEN: KDKHAY; ISSN: 0368-5098

PUBLISHER: Kogakuin Daigaku

DOCUMENT TYPE: Journal LANGUAGE: English

AB The annealing effects of Ca-La series alloys on homogenization and durability improvement were investigated, using x-ray diffraction, microg., and electrochem. measurement. The annealing was good for homogenization but showed no effect on the electrochem. durability of the La-free Ca-based alloy. When La-substituted CaNi5 type alloy was annealed at 900°C for 3 h., the phase structure changed a little, but the electrochem. durability was improved. Higher temperature and longer time annealing was effective for the electrochem. cycling durability of the Ca-La series alloys. The discharge capacity of the annealed alloys decreased because of the vaporizing of Ca during annealing for a long time.

L53 ANSWER 33 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:44265 HCAPLUS

DOCUMENT NUMBER: 124:94143

TITLE: The annealing effect on the durabilities of

La-substituted CaNi5-type alloys

AUTHOR(S): Li, Z. P.; Suda, S.

CORPORATE SOURCE: Tokyo, 192, Japan

SOURCE: Journal of Alloys and Compounds (1995), 231(1-2),

835-40

CODEN: JALCEU; ISSN: 0925-8388

PUBLISHER: Elsevier DOCUMENT TYPE: Journal LANGUAGE: English

AB The annealing effects of the Ca-La series of alloys on homogenization and durability improvement have been investigated using X-ray diffraction, microg. and electrochem. measurement. It was found that annealing has no effect on the electrochem. durability of the La-free Ca-containing alloy although it can be well homogenized. The phase structure of La-substituted CaNi5-type alloys was only changed a little by annealing the alloy at 900°C for 3h, but the electrochem. durability could be improved. A high temperature, longer anneal of the Ca-La series of alloys enabled better electrochem. cycling durability to be attained. The discharge capacity of the annealed alloys was decreased a little because of the vaporization of Ca when annealing for a long time.

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L53 ANSWER 34 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:44250 HCAPLUS

DOCUMENT NUMBER: 124:121950

TITLE: A new family of hydride electrode materials based on

CaNi5-type alloys

AUTHOR(S): Li, Z. P.; Suda, S. CORPORATE SOURCE: Tokyo, 192, Japan

SOURCE: Journal of Alloys and Compounds (1995), 231(1-2),

751-4

CODEN: JALCEU; ISSN: 0925-8388

PUBLISHER: Elsevier DOCUMENT TYPE: Journal LANGUAGE: English

AB A new family of low-cost hydride electrode materials based on CaNi5 was studied. CaNi5-series alloys showed the same charge-discharge potential as RNi5-type alloys (R = rare earth metal). A CaNi5 electrode can reach the capacity of 400 mA-h/g but only can last a few cycles in an alkaline electrolyte. The extra high capacity of CaNi5 might be brought by Ca corrosion. The CaNi5-type alloys after element substitution demonstrated a discharge capacity of 300-350 mA-h/g at a discharge current of 150 mA/g. Comparing with B-side substitution in AB5, A-side substitution is more effective on the durability improvement and keeps the discharge capacity at a high value. Rare earth and alkaline earth substitution can dramatically improve the cycling durability. This is attributed to the lattice stabilization of the alloy and size stabilization of the hydride particles as well as anti-corrosion improvement of the alloy. Zr and Ti addition in Ca0.4La0.6Ni4.7Al0.3 can improve the durability further. The A-rich non-stoichiometric alloy (AB5-type Ca-based alloy) showed higher discharge capacity but poorer durability compared with a stoichiometric compound

L53 ANSWER 35 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:44220 HCAPLUS

DOCUMENT NUMBER: 124:122029

TITLE: Effects of hydriding-dehydriding cycling on P-C-T and

electrochemical properties of La-Ca-Ni-Al alloys

AUTHOR(S): Li, Z. P.; Suda, S. CORPORATE SOURCE: Tokyo, 192, Japan

SOURCE: Journal of Alloys and Compounds (1995), 231(1-2),

594-7

CODEN: JALCEU; ISSN: 0925-8388

PUBLISHER: Elsevier
DOCUMENT TYPE: Journal
LANGUAGE: English

The effect of hydriding-dehydriding cycling on electrochem. capacity, durability, and pressure-composition-temperature (P-C-T) characteristics of Ca1-xLaxNi4.7Al0.3 alloys (x = 0, 0.2, and 0.6) was investigated at 40°. The La-substituted alloys showed a smaller electrochem. capacity degradation than CaNi5 after 40 hydriding-dehydriding cycles at 40°. Compared with uncycled alloys, La-substituted Ca-based alloys

showed better durability with the exception of La-free CaNi4.7Al0.3.

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L53 ANSWER 36 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1995:649767 HCAPLUS

DOCUMENT NUMBER:

123:95902

TITLE:

Electrochemical properties of the chemically treated

LaNi4.7Al0.3 and MmNi3.5Co0.7Al0.8

AUTHOR (S):

Yan, D. -Y.; Li, Z. -P.; Suda, S.

CORPORATE SOURCE:

Department Chemical Engineering, Kogakuin University,

Hachioji, 192, Japan

SOURCE:

Transactions of the Materials Research Society of

Japan (1994), 18B, 1217-19

CODEN: TMRJE3; ISSN: 1382-3469

DOCUMENT TYPE:

Journal English

LANGUAGE:

AB The effects of the chemical treatment by an aqueous solution which contains fluorine

(F-treatment) on the electrochem. properties of LaNi4.7Al0.3 and MmNi3.5Co0.7Al0.8 alloys was investigated. The F-treatment improved the electrochem. activation characteristics of LaNi4.7Al0.3. The treated MmNi3.5Co0.7Al0.8 improved significantly the charge discharge cycle life.

L53 ANSWER 37 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:649766 HCAPLUS

DOCUMENT NUMBER: 123:95901

TITLE: Activation of Zr0.5Ti0.5V0.75Ni1.25 electrodes by hot

alkaline solutions

AUTHOR(S): Yan, Deyi; Sandrock, Gary; Li, Zhoupeng;

Wang, Xianglong; Suda, Seijirau

CORPORATE SOURCE: Department Chemical Engineering, Kogakuin University,

Hachioji, 192, Japan

SOURCE: Transactions of the Materials Research Society of

Japan (1994), 18B, 1213-16 CODEN: TMRJE3; ISSN: 1382-3469

DOCUMENT TYPE: Journal LANGUAGE: English

AB Hot alkaline treatment of the multiphase alloy Zr0.5Ti0.5V0.75Ni1.25 results in significant surface corrosion, with at least some of the corrosion

products being soluble in the treating solution The corrosion process

generates

H that is largely absorbed by the alloy during treatment, resulting in heterogeneous particle cracking and significant electrochem. discharge capacity (CO) even before the 1st deliberate charge. The discharge capacity (C1) after the 1st deliberate charge is significantly increased by hot alkaline treatment. CO is a direct measure of alloy corrosion during the hot alkaline treatment and should be minimized in practice. C1 is a measure of electrochem. activation rate and should be maximized in practice. Another important parameter is the maximum capacity Cmax. The effects of treatment time and temperature, as well as solution (KOH vs. NaOH)

on

 ${\tt C0}$ ,  ${\tt C1}$  and  ${\tt Cmax}$  are presented, along with some long-time cyclic capacity data.

L53 ANSWER 38 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:649765 HCAPLUS

DOCUMENT NUMBER: 123:95900

TITLE: The effects of F-treatment level on the

electrochemical behavior of LaNi4.7Al0.3

AUTHOR(S): Li, Z. P.; Yan, D. Y.; Suda, S.

Department Chemical Engineering, Kogakuin University, CORPORATE SOURCE:

Hachijio, 192, Japan

Transactions of the Materials Research Society of SOURCE:

Japan (1994), 18B, 1209-12 CODEN: TMRJE3; ISSN: 1382-3469

DOCUMENT TYPE: Journal LANGUAGE: English

these results.

The F-treatment has good effects on improving the activation of LaNi4.7Al0.3 electrodes. Short-term treatments are good for higher capacities and lower charge potentials. It also can improve the electrode kinetic properties. Appropriate F-treatments can improve the galvanostatic cycling durabilities of the LaNi4.7Al0.3 electrodes without decreasing their kinetic properties. A phys. model is proposed to explain

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L53 ANSWER 39 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:487772 HCAPLUS

DOCUMENT NUMBER: 122:295255

TITLE: The fluoride protection mechanism during CaNi5

electrode cycling in the F- ion containing KOH

electrolyte

AUTHOR(S): Li, ZhouPeng; Matsuoka, Takashi; Sandrock,

Gary; Uchida, Masaki; Suda, Seijirau

CORPORATE SOURCE: Kogakuin Univ., Tokyo, 163-91, Japan

SOURCE: Kogakuin Daigaku Kenkyu Hokoku (1994), 77, 77-9

CODEN: KDKHAY; ISSN: 0368-5098

PUBLISHER: Kogakuin Daigaku

DOCUMENT TYPE: Journal LANGUAGE: Japanese

F- ion addition in KOH electrolyte can improve the cycling durability of CaNi5 electrode dramatically. Using phys. and electrochem. analyses, the durability improvement mechanism was investigated. The fluoride layers are formed on the surfaces of CaNi5 particles when discharging in F--containing KOH solution The produced CaF2 film is more protective than the Ca(OH)2 formed during cycling in KOH solution; therefore, the cycling durability of CaNi5 electrode was improved.

L53 ANSWER 40 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

1995:487771 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 123:37119

TITLE: The effect of element substitution in CaNi5 on the

durability improvement of CaNi5-type electrodes

AUTHOR (S): Li, ZhouPeng; Kanazashi, Masato; Sandrock,

Gary; Uchida, Masaki; Suda, Seijirau

CORPORATE SOURCE: Kogakuin Univ., Tokyo, 163-91, Japan

SOURCE: Kogakuin Daigaku Kenkyu Hokoku (1994), 77, 71-5

CODEN: KDKHAY; ISSN: 0368-5098

PUBLISHER: Kogakuin Daigaku

Journal DOCUMENT TYPE: Japanese LANGUAGE:

The effects of Ni substitution with Al and Co as well as Ca substitution with La in CaNi5 on the durability improvement when cycling in KOH solution were investigated. Co substitution improved the cycling durability in late cycles. Ca substitution with La also improved the cycling durability in KOH solution appreciably. However, Al substitution has no effect on the durability improvement. The structural and metallog. analyses of these alloys have been done in order to understand the durability improvement mechanism.

L53 ANSWER 41 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:469996 HCAPLUS

DOCUMENT NUMBER: 122:269997

TITLE: Electrochemical durability of Ca-based alloys

AUTHOR(S): Li, Z. P.; Suda, S.

CORPORATE SOURCE: Dep. Chem. Eng., Kogakuin Univ. Nakano, Tokyo, 192,

Japan

SOURCE: Electrochimica Acta (1995), 40(4), 467-71

CODEN: ELCAAV; ISSN: 0013-4686

PUBLISHER: Elsevier
DOCUMENT TYPE: Journal
LANGUAGE: English

AB In order to improve the cycling durability of Ca-based alloy in KOH solution as hydride electrode in Ni-H battery, we investigated the substitution effects by using Al and Co to substitute nickel as well as alkaline earth metals and rare earth elements to calcium in CaNi5. It was found that rare earth and alkaline earth substitution can significantly improve the cycling durability during electrochem. charging and discharging. Comparing with A-site substitution in AB5, B-site substitution by Al or Co shows little improvement on the durability. The structural and metallog. analyses have been done to understand the substitution effects.

L53 ANSWER 42 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:357547 HCAPLUS

DOCUMENT NUMBER: 122:138069

TITLE: The inorganic compound coating effect on the

durability of CaNi5 electrode

AUTHOR(S): Li, Z. P.; Suda, S.

CORPORATE SOURCE: Dep. Chem. Eng., Kogakuin Univ., Tokyo, 192, Japan

SOURCE: Proceedings - Electrochemical Society (1994),

94-27 (Hydrogen and Metal Hydride Batteries), 78-84

CODEN: PESODO; ISSN: 0161-6374

PUBLISHER: Electrochemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

AB F- ion addition in KOH electrolyte can improve the electrochem. cycling durability of CaNi5 electrode dramatically. The durability improvement mechanism was investigated through phys. and electrochem. analyses. Fluoride layers will be formed on the surfaces of CaNi5 particles during cycling CaNi5 electrode in F- ion containing KOH electrolyte. The formed calcium fluoride is more protective against the corrosion of strong alkaline solution than calcium hydroxide that was formed when the electrode was cycling in pure KOH electrolyte. With increasing the F- ion content in the electrolyte, the durability of CaNi5 electrode can be improved very much but the discharge capacity was decreased because of the ion conductivity loss.

L53 ANSWER 43 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:357541 HCAPLUS

DOCUMENT NUMBER: 122:138065

TITLE: The metallurgical and electrochemical investigations

of Ca-La-(NiAl)5 alloys

AUTHOR(S): Li, Z. P.; Suda, S.

CORPORATE SOURCE: Dep. Chem. Eng., Kogakuin Univ., Tokyo, 192, Japan

SOURCE: Proceedings - Electrochemical Society (1994),

94-27 (Hydrogen and Metal Hydride Batteries), 16-23

CODEN: PESODO; ISSN: 0161-6374

PUBLISHER: Electrochemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

AB The effects of lanthanum substitution for calcium in Ca1-xLaxNi4.7Al0.3 on metallog., structure and electrochem. properties. have been investigated. It was found that cored dendritic structure existed and some segregation of Ca and La occurred on solidification. Lanthanum substitution can dramatically decrease the amount of A2B7 phases, which was caused by high vapor pressure of Ca and non-equilibrium solidification of the alloy. From the results of X-ray lattice parameter measurement, the compactness of Ca-La-(NiAl)5 alloys linearly increases with increasing the La substitution amount La substitution can obviously improve the electrochem. durability of the alloy by lattice stabilization and anti-corrosion capability improvement in strong alkaline solution

L53 ANSWER 44 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1994:609064 HCAPLUS

DOCUMENT NUMBER:

121:209064

TITLE:

Rate capacity determination mechanism of F-treated

LaNi4.7Al0.3 electrode

AUTHOR (S):

Li, Zhoupeng; Yan, Deyi; Suda,

Seijirau

CORPORATE SOURCE:

Zhejiang Univ., Hangzhou, Peop. Rep. China

SOURCE:

Kogakuin Daigaku Kenkyu Hokoku (1993), 75, 113-22

CODEN: KDKHAY; ISSN: 0368-5098

DOCUMENT TYPE:

Journal English

LANGUAGE:

AGE: English
A math. model of a rate capacity determination mechanism of LaNi4.7Al0.3 as a

hydride electrode is established, based on the hydride electrode kinetics and capacity degradation mechanism. Rate cycling behavior of the F-treated

electrode was investigated. Treatment with F enhances its durability.

L53 ANSWER 45 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:609063 HCAPLUS

DOCUMENT NUMBER: 121:209063

TITLE: Effects of F treatment on durability of LaNi4.7Alo.3

electrode

AUTHOR(S): Li, Zhoupeng; Yan, Deyi; Suda,

Seijirau

CORPORATE SOURCE: Zhejiang Univ., Hangzhoud, Peop. Rep. China

SOURCE: Koqakuin Daiqaku Kenkyu Hokoku (1993), 75, 107-12

CODEN: KDKHAY; ISSN: 0368-5098

DOCUMENT TYPE: Journal LANGUAGE: English

AB Treatment of LaNi4.7Al0.3 as a hydride electrode with F to coat it with fluoride enhances its charge-discharge cycle durability. However, the

coating layer is finely cracked, when exposed to the cycles for an

extended period, which decreases capacity of the electrode.

L53 ANSWER 46 OF 46 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:609062 HCAPLUS

DOCUMENT NUMBER:

121:209062

TITLE:

Electrode kinetics of F-treated LaNi4.7Al0.3

electrodes

AUTHOR(S):

Li, Zhoupeng; Yan, Deyi; Suda,

Seijirau

CORPORATE SOURCE:

Zhejiang Univ., Hangzhou, Peop. Rep. China

SOURCE:

Kogakuin Daigaku Kenkyu Hokoku (1993), 75, 99-106

CODEN: KDKHAY; ISSN: 0368-5098

DOCUMENT TYPE:

Journal English

LANGUAGE:

An attempt was made to correlate rate capacity of LaNi4.7Alo.3 as a hydride electrode with discharge c.d. The electrode reaction is controlled by the electrochem. reaction occurring on the catalytic sites

in a high current discharge region. Treatment with F enhances kinetic

characteristics of the electrode.

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